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PROPERTIES OF ORDINARY WATER-SUBSTANCE

In all its Phases:

Water-vapor, Water, and all the Ices

N. ERNEST DORSEY, PHYSICIST National Bureau of Standards Washington, D. C.



American Chemical Society Monograph Series

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GENERAL INTRODUCTION

American Chemical Society Series of Scientific and Technologic Monographs

By arrangement with the Interallied Conference of Pure and Applied Chemistry, which met in London and Brussels in July, 1919, the American Chemical Society was to undertake the production and publication of Scientific and Technologic monographs on chemical subjects. At the same time it was agreed that the National Research Council, in cooperation with the American Chemical Society and American Physical Society, should undertake the production and publication of Critical Tables of Chemical and Physical Constants. The American Chemical Society and the National Research Council mutually agreed to care for these two fields of chemical development. The American Chemical Society named as Trustees, to make the necessary arrangements for the publication of the monographs, Charles L. Parsons, secretary of the society, Washington, D. C.; the late John E. Teeple, then treasurer of the society, New York; and Professor Gellert Alleman of Swarthmore College. The Trustees arranged for the publication of the A. C. S. series of (a) Scientific and (b) Technologic Monographs by the Chemical Catalog Company, Inc. (Reinhold Publishing Corporation, successors) of New York.

The Council, acting through the Committee on National Policy of the American Chemical Society, appointed editors (the present list of whom appears at the close of this introduction) to have charge of securing authors, and of considering critically the manuscripts submitted. The editors endeavor to select topics of current interest and authors recognized as authorities in their respective fields.

The development of knowledge in all branches of science, especially in chemistry, has been so rapid during the last fifty years, and the fields covered by this development so varied that it is difficult for any individual to keep in touch with progress in branches of science outside his own specialty. In spite of the facilities for the examination of the literature given by Chemical Abstracts and by such compendia as Beilstein's Handbuch der Organischen Chemie, Richter's Lexikon, Ostwald's Lehrbuch der Allgemeinen Chemie, Abegg's and Gmelin-Kraut's Handbuch der Anorganischen Chemie, Moissan's Traité de Chimie Minérale Générale, Friend's and Mellor's Textbooks of Inorganic Chemistry and Heilbron's Dictionary of Organic Compounds, it often takes a great deal of time to coördinate

the knowledge on a given topic. Consequently when men who have spent years in the study of important subjects are willing to coördinate their knowledge and present it in concise, readable form, they perform a service of the highest value. It was with a clear recognition of the usefulness of such work that the American Chemical Society undertook to sponsor the publication of the two series of monographs.

Two distinct purposes are served by these monographs: the first, whose fulfillment probably renders to chemists in general the most important service, is to present the knowledge available upon the chosen topic in a form intelligible to those whose activities may be along a wholly different line. Many chemists fail to realize how closely their investigations may be connected with other work which on the surface appears far afield from their own. These monographs enable such men to form closer contact with work in other lines of research. The second purpose is to promote research in the branch of science covered by the monograph, by furnishing a well-digested survey of the progress already made, and by pointing out directions in which investigation needs to be extended. To facilitate the attainment of this purpose, extended references to the literature enable anyone interested to follow up the subject in more detail. If the literature is so voluminous that a complete bibliography is impracticable, a critical selection is made of those papers which are most important.

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Preface

Origin.

This compilation was begun under the auspices of a committee of the National Bureau of Standards, the late E. W. Washburn, Chief Chemist, being chairman. Its purpose is to present either specifically or by reference all the material likely to be of interest to anyone studying the properties of ordinary water-substance, *i.e.*, that of the usual isotopic composition.*

After the general plan had been decided upon, the compiler was left entirely free to determine the details. He is greatly indebted to many for advice; but he alone is responsible for all errors of judgment in the selection of information to be included, in the form of presentation,[†] in the explanations and discussions, and in all the other details involved in the making of such a compilation.

Plan.

The plans of the committee called for (a) assembling from the *International Critical Tables* all data pertaining to the properties of the ordinary water-substance in all its phases, (b) revision and extension of those data in the light of more recent work, (c) inclusion of types of data that had been omitted from the *Critical Tables*, either through oversight or because of the nature of the plan adopted for those Tables, and (d) the arrangement of the whole so as to facilitate its use. The committee desired that the data be grouped in accordance with the several phases of the substance, and their combinations.

This general plan has been adhered to. But the term "data" has been interpreted broadly, for there is much non-numerical information that should be available to one studying the water-substance.

The units in which the numerical data are expressed are always plainly indicated, and the significance of the data is explicitly stated wherever there seems to be any danger of their being misunderstood by one not well informed in the field concerned. In some cases pertinent formulas are given or derived, and a computed quantity (e.g., disposable energy of formation) is accompanied by the basic data and the formula employed in deriving it. Most of this will seem to the expert to be very elementary and needless, but having more suitable sources of his own, he will seldom seriously consult this one for information in his own field. It is not the expert, but he who is not especially well acquainted with the field concerned, who must be considered.

^{*}A review of our knowledge of the properties of the isotope deuterium oxide (D₂O) has been published by H. C. Urey and G. K. Teal.¹

[†] Certain changes in the tabular presentations have been made by the Publisher in accordance with the style adopted for the $\Lambda.C.S.$ monographs.

¹ Urey, H. C., and Teal, G. K., Rev. Mod. Phys., 7, 34-94 (1935).

Groups of interpolation formulas (e.g., for the thermal expansion of water, Table 100) have been compared by means of skeleton tables. This has revealed some persisting errors in recognized compilations, and some oft-quoted formulas that are totally worthless. Such formulas should not appear in future compilations.

In some cases (e.g., the Verdet constant for water) an arbitrary expression suitable for computation has been set up as a norm against which to compare several independent sets of data, and by which interpolation can be readily perfermed. In no case is it claimed that such a norm represents the data. It is merely an expression that can be easily evaluated and that runs along near the data, so that deviations from it can be readily compared and studied. It is especially valuable when a wide range of the independent variable is covered by several discrete and not satisfactorily overlapping groups of observations. This procedure also has revealed some persisting errors.

Descriptive information is often given in the form of direct quotation.

Scope.

Information is given regarding the properties of pure, ordinary water-substance in all its phases—water-vapor, water, and the several ices—and regarding the phenomena and data pertaining to its synthesis and dissociation and to its transition from phase to phase; but, except as presently noted, no information is given regarding its behavior in the presence of another substance. Similar information about the water-substance as it occurs in nature has been given when readily available

The effect of the presence of air is considered, as are also the solubilities and diffusivities in water of the atmospheric and noble gases, of hydrogen, ozone, carbon monoxide, and ammonia, and the diffusion of water-vapor in air, hydrogen, and carbon dioxide, and through a few solids. All other information given concerning water and another substance is merely incidental to an understanding of the behavior of the water-substance itself.

Some types of information that might not be expected in such a compilation as this have been included. For example: The preparation of dust-free water and of monocrystals of ice, the color of water and of the sea, and the volumes of water menisci.

Period Covered.

It is hoped that no important article bearing upon the subject and appearing before 1938 has been overlooked; but only the most important of those appearing after June 30, 1937, and coming to the compiler's attention, have been considered. In accordance with the initial plan, the *International Critical Tables* has as far as possible been depended upon for information antedating January 1, 1923, and the compiler himself has searched the journals from 1922 to 1938. In many cases, data from the *International Critical Tables* have been supplemented by other early data; and in fields not covered by them, the compiler has tried to consult all the

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significant reports, early as well as recent. He fully realizes that he has only partially succeeded in this attempt, and will be glad to have important omissions brought to his attention.

Acknowledgments.

It is a pleasure to the compiler to acknowledge his great indebtedness to the many who have assisted him in divers ways: to the *International Critical Tables* and its several experts, and to the National Academy of Sciences for their gracious permission to use the data and other information published in those tables; to Messrs. Friedr. Vieweg & Sohn, Braunschwieg, Germany, for their kind permission to use Tables 41 and 42 of the Warmetabellen prepared by Holborn, Scheel, and Henning, and published in 1919; to the various investigators who have assisted him by correspondence regarding their own work; to his fellow associates, members of every interested division of the National Bureau of Standards, for information, advice, and criticism; and especially to Dr. Lyman J. Briggs, Director of that Bureau, for his unfailing patience and encouragement, without which the work could not have been done.

N. ERNEST DORSEY.

National Bureau of Standards, Washington, D. C. September, 1938.

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Introduction

General information that will facilitate the use of this compilation is given in this Introduction. It is classified under two heads: (1) arrangement and documentation, and (2) symbols, units, and equivalents. Information regarding the origin, plan, and scope of this volume, and the period covered by it, will be found in the Preface.

1. Arrangement and Documentation

Arrangement.

As may be seen in the table of contents, information regarding the water-substance has been assembled in five broad groups: (I) Synthesis and dissociation; (II) Single-phase systems, subdivided into (IIa) Water-vapor, (IIb) Water, (IIc) Ice; (III) Multiple-phase systems; (IV) Phase transition; and (V) Miscellanea. Each of these is subdivided into smaller units, devoted to a closely related group of data or phenomena.

As far as possible, the subgroups are arranged according to the nature of the phenomena involved, and in the following order: atomic or molecular, mechanical, acoustic, thermal, optical, electrical and magnetic.

Everything that either involves the presence of a second phase (e.g., surface tension) or requires the presence of a second phase in order to insure the existence of the assumed condition (e.g., pressure of saturated vapor) has been placed in Group III (Multiple-phase systems), excepting such (e.g., latent heat) as have to do with the act of transition from one phase to another. The last form the contents of Group IV (Phase transition). In Group V (Miscellanea) have been placed certain odd bits of information that do not fit in well elsewhere, and a brief note on interpolation.

Documentation.

All data and descriptive information, and most of the formulas and statements regarding theories, are accompanied by references to the sources from which they have been obtained or on which they rest. In many cases references to secondary sources that have come to the compiler's attention and seem to be of importance or interest to one seriously studying the subject are given also, although not otherwise used in the compilation.

With the exception of references taken from the *International Critical Tables*, and a few others plainly indicated as secondary, each book and article cited has been examined by the compiler with reference to the information accredited to it; and so far as it was practical, the limiting pages have been determined for the references from the *Critical Tables* also; but

of the papers covered by those references, only those pertaining to fields that had to be reworked have been studied by the compiler.

If the results of an investigation have been published more than once, whether in whole or in part, in abstract, as advance notice, or as a correction, a reference to each such publication that has come to the compiler's attention is given, except that no reference is given to the Bulletin of the American Physical Society, the abstracts in that being contained in the Physical Review also, to which reference is made.

The relations between the several articles as regards extent and content, but not time of publication, are indicated by means of the following symbols used for separating the references, each referring primarily to the two between which it appears: = means that the articles differ in no essential feature other, perhaps, than language; — means that one is a slightly revised copy of the other, the changes may or may not be of importance; \rightarrow and \leftarrow mean that the one at the head of the arrow is a shorter report (abstract, review, etc.) than that at the other end. The shorter is always at the head of the arrow, whatever its date.

In all cases, the information to which such a string of references relates has been derived from the first of the string, unless the contrary is explicitly stated.

Occasionally it has seemed desirable to refer to a particular page or illustration, or to give the characteristic designation of the article as a separate unit in some recognized series, or to indicate whether the article is an abstract (A), a letter to the editor (L), or a review (R). In such cases, this parenthetical information is enclosed in parentheses and placed after the final page number of the article, the letters A, L, and R being used as here indicated.

The year given in a reference is generally that given on the title page of the volume containing the paper referred to. If the title page indicates that the volume covers portions of two or more years, then the last of those years is the one generally given. Some such rigid rule is necessary if simplicity is to be secured, for the covers of some reprints have a date that is incompatible with the dates on the title page of the volume containing the article in question. For example, the title page of volume 70 of the Proceedings of the American Academy of Arts and Sciences states that it covers from May 1934 to May 1935, and it carries 1936 as the date of publication. To the nine papers that it contains are attached the following dates:

Paper	Received	Presented	Cover date
1	October 1, 1934	October 10, 1934	March, 1935
2	October 8, 1934	February 13, 1935	March, 1935
3	December 14, 1934	December 12, 1934	April, 1935
4	February 4, 1935	February 13, 1935	May, 1935
5	May 9, 1935	March 13, 1935	August, 1935
6	August 6, 1935	March 13, 1935	December, 1935
7	October 15, 1935	October 9, 1935	December, 1935
8	December 3, 1935	December 11, 1935	February, 1936
9	December 12, 1935	December 11, 1935	February, 1936

A reference to any paper in that volume will in the present compilation be dated 1935.

2. Symbols, Units, and Equivalents

To facilitate the use of this volume, the symbols used in any major section have usually been defined therein, the exceptions being mainly the well-standardized symbols, including those of the units. These and a few others that may be unfamiliar to some have been assembled in Table 1, where certain of them are defined, and some numerical values and conversion factors are given. Of the symbols for units, however, only the simpler are included, from which the more complicated may be constructed in the well-known manner described in a following paragraph. Many of the symbols, other than those for units, are occasionally used with other meanings, but the context, and especially the definitions in the accompanying text, will enable the user to determine the proper interpretation in each case.

Italics.

In accordance with the established custom of the English language, English letters used out of character, that is, in senses different from those ascribed to them in usual English script, are printed in distinctive type, in italics if the context is roman, and conversely. Those used in character may also be printed in distinctive type for very special purposes. Exceptions occur, which are usually in the direction that leads to conformity with other members of the group of related symbols.

Of the abbreviating symbols used in this compilation, true abbreviations and the symbols for the units of measure, for the chemical elements, and for the names of mathematical functions are printed in roman; most of the others are printed in italics, the letters composing them being obviously used out of character.

Symbols of Units.

Symbols of units are printed in Greek or Roman letters, never in italics, and generally do not end with a period, the principal exceptions to the last being the symbols for the British units (in., ft., lb., etc.), and the period is not always used with them. In writing the synthetic symbol for a derived unit, the symbols of the units that form a product are separated by a period without additional space (g·cm); those that form a ratio or a quotient are either combined as a fraction $\left(\frac{cm}{sec}\right)$ or separated by the shilling mark (cm/sec) or the symbols in the divisor are written with negative exponents and separated from those of the dividend by a period without additional space (cm·sec⁻¹, g·cm·sec⁻²). In all such cases any period that might otherwise form part of the symbol of an individual unit is omitted, thus avoiding a duplication of the period in the interior; the final period

is unnecessary, as the resulting combination is of a type that is never used except as a symbol for a unit. The periods indicating multiplication are placed above the line.

The elements of the symbol of a unit designated by a compound name in which the elements of the compound are not to be understood as being combined by processes analogous to multiplication and division, are connected by a hyphen (ft-c = foot-candle, mm-Hg = millimeter of mercury).

In general, the same form of symbol (the singular) is used whatever the magnitude of the numeral to which it is attached. When convenient, the symbol is preceded, without spacing, by an integral power of ten (positive or negative), to indicate that the unit used is so related to that designated by the unmodified symbol ($2.9986 \ 10^{10}$ cm/sec = 2.9986×10^{10} cm/sec).

Prefixes.

The following metric prefix symbols and prefixes are used, each indicating that the ratio of the unit to that corresponding to the symbol or name to which the prefix is attached is that shown by the corresponding number, e.g., 1 microgram = 0.000 001 g, 1 kilogram = 1000 g.

 μ - = micro- = 0.000 001, m- = milli- = 0.001, c- = centi- = 0.01, k- = kilo- = 1000, mega- = 1000 000. There is no generally accepted single-character symbol for mega. (A capital M is sometimes used, but that may be misread for the commonly used Roman numeral for 1000; the Roman symbol (\overline{M}) for a million seems to be more appropriate.) Each of these symbols has always the significance here given; m- never stands for either micro- or mega-, but solely for milli-. The final vowel of the prefix is dropped from mega- and from micro- when the next letter is a vowel.

These prefixes are combined, one to another, as need be, the one nearer to unity preceding the other ($m\mu$ - millimicro- = 10^{-9} ; kmega- = kilomega- = 10^{9}) never the other way round. Either singly or combined they may be directly attached to any unit of, or based upon, the metric system.

Systems of Units.

The normal system of units based upon the centimeter, the gram (unit of mass), and the second, and commonly designated as the cgs system, is generally used; but the practical absolute electrical units, *i.e.*, ohm = 10^9 cgsm, ampere = 0.1 cgsm, volt = 10^8 cgsm, joule = 10^7 cgsm = 10^7 ergs, watt = 10^7 erg/sec, and their international counterparts are also used; and occasionally the British units (ft., lb., etc.).

The cgs electrostatic system of units is denoted by the symbol cgse; the electromagnetic, by cgsm. The same three symbols may be used to denote the appropriate unit in the corresponding system, whatever the nature of that unit, as cgsm is used in the preceding paragraph.

As the names of the several international electrical units are the same as those of their counterparts in the practical absolute system, it is occasionally necessary to distinguish between the two. In such cases, the qualifier "(Int.)" accompanies the symbol for the unit when that must be interpreted as the international unit. If this qualifier does not appear, the unit is the absolute one, or the number to which the symbol is attached is not known with sufficient accuracy to justify making a distinction between the two. It is only with respect to the joule that it is ever necessary to make the distinction in this compilation.

It must be remembered that the international joule used in experimental work is defined in terms of the international ohm and the international volt, and that the certification of resistances and of standard cells is always in terms of the concrete standards of the certifying laboratory; those standards define the international units for that laboratory. The amount of energy corresponding to the international joule as defined by such laboratory units of resistance and voltage has varied from laboratory to laboratory, and from time to time, the variation steadily decreasing as the concrete standards have been improved in permanence and in reproducibility. even as late as 1931 it was very significant. At that time the international joule of Great Britain, as defined by the ohm and the volt, was greater than those of Germany, France, and this Bureau, the differences being, respectively, 1.91, 1.98, and 0.62 parts in 10 000 (hundredths of a per cent).1 Since that comparison, some of the national laboratories have revised the values assigned to their concrete standards; and now (1938) the discrepancies between the values of the international joules of the several countries amount to no more than 3 or 4 parts in 100 000 (3 or 4 thousandths of a per cent). It is confidently expected that with the present arrangements for systematic intercomparisons and the use of the better standards now available, the discrepancies will be kept continuously well below those existing in 1931. At present (1938), the international joule as defined by the standards of the National Bureau of Standards lies between 1,0002 and 1.0003 absolute joules, and is probably nearer the lower value.

From this discussion it is obvious that when one has to do with an accuracy of a hundredth of a per cent, or higher, it is impossible to translate observations expressed in international joules into absolute joules without loss of accuracy, unless one knows what particular international joule was used, and how it is related to those for which comparisons with the absolute joule have been made. This is true irrespective of the accuracy of the absolute measurements.

For this reason the compiler has not attempted to convert reported data from one of these joules to the other, unless the author himself has given the conversion factor that he believed to apply.

¹ Vinal, G. W., Bur. Stand. J. Res., 8, 729-749 (RP448) (1932).

Conversion Factors.

Such conversion factors as seem appropriate are given immediately above the numerical data in each table, in the following form: Unit of $P = 1 \text{ kg*/cm}^2 = 0.967841 \text{ atm} = 735.559 \text{ mm-Hg} = 0.980665 \text{ bar}$. This indicates that a number, say a, standing in the P-column, represents a pressure of a kg*/cm², of 0.967841a atm, of 735.559a mm-Hg, of 0.980665abars, all of which pressures are equal, one to another.

Table 1.—Symbols, Units, and Equivalents

With a few exceptions, this table contains only the simple units of measure appearing frequently in this compilation, certain well-standardized symbols that are not always defined in the accompanying text, and some symbols and names that may be unfamiliar to the user, and none of the well-known mathematical symbols. Some of these symbols, other than those for the units, are sometimes used in ways not indicated here; those uses are sufficiently explained where they occur. The synthetic symbols for the derived units of measure are formed from the simple ones in the well-known way already described.

In the first column are given alphabetically the abbreviating symbols and the names of those units for which there are no such symbols. In the second are given the names of the units or other quantities corresponding to the symbols, and the definitions and equivalents of the units, or as much of this as seems desirable.

Greek symbols are given at the end of the table.

Α Angstrom, a unit of length; $1A = 10^{-8}$ cm = 0.1 m μ .

A A pressure equal to 1 atm; its numerical value depends upon the unit of pressure.

atm Normal atmosphere, a unit of pressure; 1 atm = 1.01325 mega $dynes/cm^2 = 1.03323 \text{ kg*/cm}^2 = 1.01325 \text{ bars.}$ Note: In Germany, "at" is frequently used to denote a pressure of $1 \text{ kg*/cm}^2 = 0.96784 \text{ atm.}$

Bar, a unit of pressure; $1 \text{ bar} = 1 \text{ megadyne/cm}^2$. bar

°C Degree centigrade. The degree interval on the scale of the centigrade thermometer, on which the normal melting point of ice is called 0, and the normal boiling point of water is called 100. Unless something else is clearly indicated, the intervals are to be counted from that zero.

Centi-, a prefix meaning 1/100. c-

Specific heat at constant pressure. c_p

Specific heat at constant volume. C ,,

Gram calorie, a unit of heat. Unless another value is specified, cal

it is assumed that 1 cal = 4.185 joules.

15°-calorie. Similarly for cal₂₀, cal_m. See Section 33. cal₁₅

Table 1—(Continued)

cgs A symbol used to designate either the system of normal units based up the centimeter, the gram (unit of mass), and the second, or a unit of that system.

cgse A symbol for any unit of the cgs electrostatic system, and for the system itself.

cgsm A symbol analogous to cgse, referring to the cgs electromagnetic units and system.

cm Centimeter, a unit of length; 1 cm = 0.01 m = 0.032808 ft.

cm-Hg Centimeter of mercury, a unit of pressure; 1 cm-Hg = 13.3322 kilodynes/cm² = 13.3322 millibars. By definition, 1 cm-Hg is the pressure exerted by a vertical column of mercury 1 cm long, at a place where the acceleration of gravity is 980.665 cm/sec², and when the density of the mercury is 13.5951 g/cm³ and either the free surface of the column is flat or a proper correction has been made for the effect of its curvature.

Density; seldom used. Also, derivative.

d

е

dyne The cgs unit of force; the force that will give to a mass of one gram an acceleration of 1 cm/sec².

E Internal or intrinsic energy of a substance or system.

The number 2.71828..., the base of the natural system of logarithms.

erg The cgs unit of work and energy; the work done by a constant force of one dyne while the point of application of the force moves one centimeter in the direction in which the force acts; 1 erg = 1 cm·dyne.

OF Degree Fahrenheit. The degree interval on the scale of the Fahrenheit thermometer, on which the normal melting point of ice is marked 32, and the normal boiling point of water is marked 212. Unless something else is clearly indicated, the intervals are to be counted from the origin defined by these numbers.

ft. Foot, unit of length; 1 ft. = 30.480 cm.

The acceleration of gravity. By international agreement all data involving the value of g are to be reduced to the basis of g = 980.665 cm/sec² = 32.1740 ft/sec², which is called the "normal" value of g.

g Gram, a unit of mass; 1 g = 0.0022046 lb (avdp.)

g* Gram weight, a unit of force; the weight of a mass of 1 g at a place where g = 980.665 cm/sec²; 1 g* = 980.665 dynes.

gfw Gram formula weight, a unit of mass; as many grams as there

Table 1—(Continued)

are units in the formula weight. In any specific case the pertinent formula should be clearly indicated.

gfw- H_2O Gram formula weight of H_2O ; 1 gfw- H_2O = 18.0154 grams of the water-substance.

g-mole Gram mole, a unit of mass; as many grams as there are units in the molecular weight.

H Enthalpy; heat content; total heat. $\Delta H = \Delta (E + pv)$, where Δ means "the increase of," p = pressure, v = volume, E = intrinsic energy. In the case of the water-substance, H is commonly used to denote the value of ΔH in going from saturated water (liquid) at 0 °C to the indicated state.

h Planck's constant of action; $10^{27} h = 6.56 \text{ erg-sec.}$

in. Inch, a unit of length; 1 in. = 2.5400 cm.

(Int.) International. (Int.) accompanies the symbols for units belonging to the international electrical system.

j Joule, a unit of energy; $1 \text{ j} = 10^7 \text{ ergs} = 10 \text{ megergs}$.

°K Degree Kelvin. The degree interval on that thermodynamic scale of temperature which has 100 degrees between 0 °C and 100 °C. Unless something else is clearly indicated, the intervals are to be counted from the absolute zero, which in this compilation is assumed to be 273.1 °K below 0 °C, unless another value is definitely specified, cf. Table 266.

k Boltzmann's constant; molecular gas-constant. $k \equiv R/N = 1.372 \, 10^{-16} \, \text{erg/}^{\circ} \text{K}$ per molecule.

k- Kilo-, a prefix meaning 1000.

kc Kilocycle = 1000 cycles. kg Kilogram; 1 kg = 1000 g.

kg* Kilogram weight; 1 kg* = 1000 g* = 980665 dynes.

kij Kilojoule; 1 kj = 1000 j. km Kilometer; 1 km = 1000 m.

Liter, a unit of volume; $1 1 = 1000.027 \text{ cm}^3$. By Pound; a unit of mass; 1 lb = 453.59243 g.

lb* Pound weight, a unit of force. The weight of a mass of 1 lb at a place where g = 980.665 cm/sec²; 1 lb* = 0.44482 megadyne.

(lb* in²)_L A pressure of one pound per square inch at a place (London) where the acceleration of gravity is 981.16 cm/sec²; 1 (lb*/in²)_L = 1.000505 lb*/in² = 68.982 millibars.

log_e Logarithm to the base e; natural logarithm. $Log_{e}x = 2.302585 log_{10}x.$

log₁₀ Logarithm to the base 10; common logarithm.

Table 1—(Continued)

Molecular weight. For H_2O , M = 18.0154.

m Mass.

m- Milli-, a prefix meaning 1/1000.

m Meter, a unit of length; 1 m = 100 cm = 3.2808 ft.

mega- A prefix meaning 1 000 000. mg Milligram; 1 mg = 0.001 g.

mg* Milligram weight, a unit of force; 1 mg* = 0.001 g*.

micro- A prefix meaning 1/1 000 000.

micron A unit of length (see μ).

ml Milliliter; 1 ml = $0.001 \ 1 = 1.000 \ 0.0027 \ cm^3$. mm Millimeter; 1 mm = $0.001 \ m = 0.1 \ cm$.

mm-Hg Millimeter of mercury, a unit of pressure (see cm-Hg);

1 mm-Hg = 0.1 cm-Hg = 1.33322 millibars.

ms Millisecond; 1 ms = 0.001 sec.

mμ- Millimicro-, a prefix meaning 1/1 000 000 000.

m μ Millimicron; 1 m μ = 0.001 μ = 10⁻⁹ m = 10⁻⁷ cm = 10A.

N Avogadro's number; the number of molecules per gram-mole. $N = 6.061 \times 10^{23}$.

P, p Pressure.

p Poise; see below.

poise The cgs unit of viscosity. It is the viscosity of a liquid which, when streaming lamellarly, exerts upon one side of an internal layer parallel to the lamellas a drag of 1 dyne/cm² in the direction of the velocity v when the value of dv/dx at that side of the layer is 1 cm/sec per cm, dx being an element of the normal drawn outward from that side of the layer;

1 poise = 1 g/cm·sec = 1.0197 mg*-sec/cm² = 1 dyne·sec/cm².

R The gas-constant. $R = 8.315 \text{ j/g-mole} \cdot \text{°K}$.

radian A unit of angle. The angle of which the arc is equal to the radius; 1 radian = 57° 17' 44.8".

r.m.s. Root-mean-square. The square root of the mean of the squares of the individual values of the quantity indicated.

sat Saturated; at saturation. Used chiefly as a subscript.

steradian A unit of solid angle. The solid angle subtended at the center by a spherical surface equal in area to the square of the radius of the sphere. The solid angle subtended at its center by a hemisphere is 2π steradians.

Temperature, on the thermodynamic scale.

t Temperature, on the centigrade scale. Occasionally, time.

Table 1—(Continued)

ton A unit of mass; 1 (short) ton = 2000 lb, 1 (long) ton = 2240 lb.

ton* Ton weight, a unit of force; 1 (short) ton* = 2000 lb*, 1 (long) ton = 2240 lb*.

v Volume. Velocity.

 v^* Specific volume, the volume of a unit of mass of the substance.

W Disposable energy. See Section 6.

 γ Ratio of the principal specific heats of a substance; $\gamma = c_p/c_v$.

 Δ A deviation or difference. $\Delta x = \text{an increase in } x$.

δ A deviation or difference.

ε Dielectric constant.

 η Viscosity.

κ Magnetic susceptibility. Electrical conductivity.

 Λ Equivalent conductivity (electrical).

λ Wave-length.

μ Magnetic permeability. Joule-Thomson coefficient. Moment of a dipole. Coefficient of absorption.

 μ Micron, a unit of length; $1 \mu = 10^{-6} \text{ m} = 0.001 \text{ mm} = 10\,000\text{A}$.

 μ - Micro-, a prefix meaning 1/1~000~000.

 μ -Hg Micron of mercury, a unit of pressure; 1 μ -Hg = 0.001 mm-Hg.

ν Wave-number; ν = 1/λ. Number. Frequency.

Pi, the ratio of the circumference of a circle to its diameter; $\pi = 3.14159...$

ρ Density. Depolarization factor.

σ Density (seldom used).τ Time. Transmissivity.

χ Specific susceptibility (magnetic).

Cycle.

~/sec Cycle per second.

I. Synthesis and Dissociation

IA. SYNTHESIS

3. Union of Hydrogen and Oxygen

The chemical reactions that occur in mixtures of hydrogen and oxygen, with or without the admixture of another gas; the way they vary with the temperature, pressure, illumination, and composition of the mixture; the ignition temperatures and explosion limits of such mixtures, and the way these vary with the size, form, and material of the containing vessel; and all the other various phenomena associated with the reactions that occur in such mixtures:—all these lie beyond the scope of this compilation, which in the field of the synthesis of water is limited to the stoichiometric composition of water, the heat of formation, and the maximum work that can be obtained from the reaction when it is carried out at constant pressure.

Those desiring information regarding the chemical reactions themselves, the attendant phenomena, and the way they vary with the conditions are referred to the compilations by W. A. Bone and D. T. A. Townend,¹ A. Skrabal,² and C. Winther.³ Those desiring more recent data, conclusions, and inferences can obtain from the papers listed in the accompanying notes ⁴⁻²⁸ a general idea of the present status of the subject, and references

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<sup>1</sup> Bone, W. A., and Townend, D. T. A., Int. Crit. Tables, 2, 172-195 (1927).
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² Skrabal, A., Idem, 7, 113-152 (1930).

³ Winther, C., Idem, 7, 159-173 (1930).

⁴ Alyea, II. N., and Haber, F., Z. physik. Chem., (B), 10, 193-204 (1930).

⁵ Andreew, K. K., and Chariton, J. B., Trans. Faraday Soc., 31, 797-804 (1935).

⁶ Bestchastny, A. L., ct. al., Physik. Z. Sowj., 5, 562-579 (1934).

⁷ Breton, J., and Laffitte, P., Compt. rend., 202, 316-318 (1936).

⁸ Chapman, D. L., and Reynolds, P. W., Proc. Roy. Soc. (London) (A), 156, 284-306 (1936).

ⁿ David, W. T., Phil. Mag. (7), 20, 65-68 (1935). Nature, 138, 930 (L) (1936); 139, 67-68 (L) (1937).

¹⁰ Drop, J., Rec. Trav. Chim. Pays-Bas, 54, - (4), 16, 671-679 (1935).

¹¹ van Heiningen, J., Rec. Trav. Chim. Pays-Bas, 55, 65-75, 85-100 (1936).

¹² Hinshelwood, C. N., ct. al., Proc. Roy. Soc. London, (A), 118, 170-183 (1929); 122, 610-621 (1929); 124, 219-227 (1929); 130, 640-654 (1931); 134, 1-7 (1931); 138, 311-317 (1932); 141, 29-40 (1933). Trans. Faraday Soc., 28, 184-191 (1932). Nature, 131, 361-362 (1933). Z. Elektroch., 42, 445-449 (1936).

¹⁸ Hinshelwood, C. N., and Williamson, A. T., "The reaction between hydrogen and oxygen," Oxford University Press, 1934.

¹⁴ Jost, W., Z. Elektroch., 41, 183-194, 232-250 (1935); 42, 461-467 (1936).

¹⁵ Lewis, B., and von Elbe, G., J. Chem'l Phys., 2, 537-546 (1934); J. Am. Chem. Soc., 59, 656-662 (1937); 59, 970-975 (1937).

¹⁰ Lindeijer, E. W., Rec. Trav. Chim. Pays-Bas, 56, 97-104, 105-118 (1937).

¹⁷ Maas, J. H., and Ewing, C., J. Phys'l Chem., 37, 13-15 (1933).

¹⁸ Malinowski, A. E., and Skrynnikow, K. A., Physik. Z. Sowj., 7, 43-48 (1935).

¹⁰ Miyanishi, M., Sci. Papers Inst. Phys. Chem. Res. (Tokyo), 26, 70-76 (1935); 27, 47-51, 52-58 (1935).

²⁰ Mole, G., Proc. Phys. Soc. (London), 48, 857-864 (1936).

²¹ Poljakow, M. W., et. al., Acta Physicochim. URSS, 1, 551-553, 817-820, 821-832 (1935); 2, 211-214, 397-400 (1935).

to other pertinent publications. The list does not pretend to include all the important papers, nor even all the most important ones, but only those which happen to be readily at hand. In certain cases of multiple authorship the references have been listed under the name of a single author, even though that name appears in some of the cases as junior author.

THE COMPOSITION OF WATER

A review of the best work that has been done on the stoichiometric composition of water has been written by J. R. Partington.²⁹ He concluded that the ratio of the combining volumes at 0 °C and a pressure of 760 mm-Hg is $O_2/H_2 = 1/2.00288$.

If the atomic weight of II is 1 0077,30 and that of O is 16.0000, the molecular weight of H_2O is M = 18.0154.

Natural water contains 1 D₂ to about 6500 H₂ 31

HEAT OF FORMATION OF H₀O

By the heat of formation $(Q_{TP})_{\phi}$ of a certain phase (ϕ) of a substance at pressure P and absolute temperature T is meant the amount of heat evolved when one gram-formula-weight (gfw) of ϕ at P and T is formed from its elements at T and the total pressure P. It is the decrease in the enthalpy (H = E + pv) at fixed T and P per gfw of the substance formed.

Its value for any P and T within the domain in which the phase ϕ can exist can be determined from its value for one such pair of values by means of relation (1) in which δ may be computed by means of any one of the relations (2a), (2b), etc.

$$(Q_{T,P})_{\phi} = (Q_{T,P})_{\phi} - \delta \tag{1}$$

$$\delta = \int_{T'}^{T} (\Delta C)_{P} dT + [P(\Delta v)_{P} - P'(\Delta v)_{P'}]_{T} + [(\Delta E)_{P} - (\Delta E)_{P'}]_{T}$$
(2a)

$$= \int_{T'}^{T} (\Delta C)_{P} dT + [P(\Delta v)_{P} - P'(\Delta v)_{P}]_{I} + [(\Delta E)_{P} - (\Delta E)_{P'}]_{T'} (2b)$$

$$= \int_{P'}^{T} (\Delta C)_{P} dT - \int_{P'}^{P} [\Delta(\mu C)]_{T} dp$$
 (2c)

- -2 Prettre, M, and I iffitte, P, Compt rend, 187, 763 765 (1928), 188, 397-399 (1929)
- ²⁴ Prettre, M., Compt. rend. 196, 1891-1893 (1933), 201, 962 964 (1935), 204, 1734 1736 (1937) ²⁴ Rodebush, W. II., et al., J. Am. Chem. Soc., 59, 1924 1931 (1937), J. Phys'l Chem., 41, 283 291 (1937).
- Semenoff, N. 7 Physik, 48, 571-582 (1928); 7 physik Chem. (B), 2, 161-168, 169-180 (1929). (B), 28, 43 53, 54 64 (1935). Chem Rev., 6, 347 379 (1929)
 Semenova, N. Acta Physicachim URSS 6, 25-42 (1937).
 - 27 Sokolik, A., and Shtsholkin, K., Acta Physicochim. URSS 1, 311 317 (1934)
 - 28 Taurin, P, Compt rend 196, 1605 1607 (1933), 197, 1046 1049 (1933).
 - 29 Partington, J. R, "The Composition of Water," 1928
 - 80 Int Crit Tables, 7, 44 (1926)
- ³¹ Hall, N. F. and Jones, T. O. I. Am. Chem. Soc., 58, 1915-1919 (1936), and Gabbard, J. L., and Dole, M., Idem, 59, 181 185 (1937).

$$= \int_{T'}^{T} (\Delta C)_P dT - \int_{P'}^{P} [\Delta'(\mu C)]_{T'} dP$$
 (2d)

Here, ΔC , Δv , and ΔE denote, respectively, the increase in the specific heat at constant pressure, in the specific volume, and in the internal energy per unit mass, when the elements taken in the right proportions combine to form the substance, the pressure and temperature being those indicated by the subscripts, and the unit of mass being in all cases that of one gfw of the substance formed; the Joule-Thomson coefficient μ is $(\delta T/\delta p)_a$; p is a variable pressure; the a indicates that the change is adiabatic; and $\Delta(\mu C)$ is the excess of μC for ϕ over that for the equivalent mixture of the uncombined elements.

If ϕ and all the elements involved may be regarded as ideal gases in the domain considered, then $(\Delta C)_P = (\Delta C)_{P'}$, $\mu = 0$, and all except the first

term in each expression for
$$\delta$$
 is zero, giving $\delta = \int_{T'}^{T} (\Delta C)_{P} dT$. Also, if ϕ is

either a liquid or a solid, and is formed from gaseous elements, its volume will usually be negligible as compared with that of the gases from which it is formed, and then δ again reduces to its first term if the gases are ideal and the changes in the volume and in the internal energy with the pressure are negligible for ϕ .

This simplification is frequently assumed, but it is valid only under the condition just stated. In many cases, especially when the pressure approaches or equals that of saturation of one of the substances, data are not available for determining the error so introduced. For water-vapor, there are no data for μ in that region, and those in the region of superheat indicate that μ increases rapidly as saturation is approached.

If P and T are such associated values that the phases ϕ and ϕ' are in equilibrium, and if L_{TP} is the latent (absorbed) heat per gfw on passing from ϕ' to ϕ at P and T, then $(Q_{TP})_{\phi} = (Q_{TP})_{\phi'} - L_{TP}$.

From these relations it is possible to compute the heat of formation of any phase of a substance for any allowable P and T if the value of (Q_{TP}) for any one phase and one set of values of P and T is known, together with the values of the required auxiliary quantities.³²

Whenever the data in the following paragraphs are expressed in international joules and are of such a precision as to justify a distinction between that unit and the absolute joule, the symbol "(Int.)" will precede the symbol for the unit. This symbol does not appear when the precision of the data is too low to justify any distinction between the units, or when the data are expressed in terms of the absolute unit; in either case, the unqualified symbol for the unit may properly be interpreted as indicating the absolute unit.

²⁵ For a review of the available data, see Bichowsky, F. R., and Rossini, F. D., "The Thermochemistry of the Chemical Substances," New York, Reinhold Publishing Corp., 1936.

Water-vapor: Heat of Formation.

If the heat of formation of water may be regarded as independent of the pressure when that does not exceed one atmosphere, as is essentially true, then the heat of formation of water-vapor at 25 °C and the pressure of water-vapor saturated at that temperature may be derived from Rossini's value (p. 17) for the heat of formation of water (285.775 (Int.) kj/gfw = 285.890 kj/gfw at 25 °C and one atmosphere) and the latent heat of vaporization at 25 °C, 43.939 (Int.) kj/gfw = 43.956 kj/gfw.³³ It is

$$(Q_{298.1,23.76 \text{ mm}})_g = 241.84 \text{ (Int.) kj/gfw} = 241.93 \text{ kj/gfw-H}_2\text{O}$$

= 13.424 (Int.) kj/g = 13.429 kj/g

With this value those given in *International Critical Tables* (242.0 at 18 °C ³⁴ and 241.8 at 25 °C ³⁵) agree remarkably well.

Bichowsky based his value on the sources from which he derived his value for the heat of formation of water (p. 17), together with the sources given below,³⁶⁻⁴¹ which treat of the heat of vaporization. (From the way in which these references are given in the *Critical Tables* one might infer that the data in them were used alone, instead of in conjunction with those contained in the immediately preceding references for the formation of water.)

Randall's value was not derived from the data contained in the references heading the table in which it appears, but seems to have been arrived at in the following manner: From the work of A. Schuller and V. Wartha, 42 of C. von Than, 43 and of J. Thomsen, 44 G. N. Lewis 45 deduced the value 68470 cal/gfw for the heat of formation of water at 0 °C and 1 atm, and from that the value 68270 cal/gfw at 25 °C. From the work of T. W. Richards and J. H. Mathews, 40 and of A. W. Smith, 41 G. N. Lewis and M. Randall 46, p. 477 chose 540.0 cal/g or 9730 cal/gfw as the value of the latent heat of vaporization of water at 0 °C, and from that obtained 10450 cal/gfw at 25 °C. Whence the heat of formation of water-vapor is 57820 cal/gfw at 25 °C and 1 atm.

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** Osborne, N. S., Stimson, H. F., and Fiock, E. F., Bur. Stand. J. Res., 5, 411-480 (RP209) (1930).

** Bichowsky, F. R., Int. Crit. Tables, 5, 176 (1928).

** Randall, M., Idem, 7, 231 (1930).

** Carlton-Sutton, T., Proc. Roy. Soc. (London) (A), 93, 155-176 (1917).

** Griffiths, E. H., Phil. Trans. (A), 186, 261-341 (1895).

** Henning, F., Ann. d. Physik (4), 21, 849-878 (1906); (4), 29, 441-465 (1909); (4), 58, 759-760 (1919).

** Holborn, L., Scheel, K., and Henning, F., "Warmetabellen," Vieweg, Braunschweig, 1919.

** Am. Chem. Soc., 33, 863-888 (1911).

** Smith, A. W., Phys. Rev., 33, 173-183 (1911).
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⁴² Schuller, A., and Wartha, V., Ann. d. Physik (Wied.), 2, 359-383 (1877).

⁴⁸ von Than, C., Idem, 13, 84-105 (1881).

⁴⁴ Thomsen, J., 1dem, (Pogg), 148, 368-404 (1873).

⁴⁵ Lewis, G. N., J. Am. Chem. Soc., 28, 1380-1395 (1906).

⁴⁰ Lewis, G. N., and Randall, M., "Thermodynamics and the Free Energy of Chemical Substances," New York, McGraw-Hill Book Co., 1923.

Lewis and Randall 46, p. 59 state that this calorie is equivalent to 4.182 × 10⁷ ergs, and Randall ⁴⁷ states that it is equivalent to 4.182 joules. Each statement implies that this calorie is to be regarded as equivalent to 4.182 absolute joules; and that interpretation is borne out by A. W. Smith's article 41 to which they refer. They have essentially followed Smith, who averaged the result obtained from the absolute (mechanical) determination by O. Reynolds and W. H. Moorby 48 with that from the electrical determination by H. T. Barnes 49 as converted to the basis of the 1911 international joule. This procedure tacitly assumes that the international joule of 1911 is essentially identical with the absolute joule, and that the average so obtained may be regarded, as it is by Smith, as the value of the calorie in terms of the absolute joule.

If the value 4.182 given by Lewis and Randall is used as the conversion factor, 57280 cal/gfw becomes 241.8 kj/gfw, the value given by Randall (see p. 14). But Lewis and Randall 45, p. 59 state that their unit is the 15°-calorie, which is actually close to 4.185 joules, making 57280 cal/gfw = 242.0 kj/gfw. Which of these two is to be preferred, and what importance is to be attached to the digit following the decimal point can be determined only by a detailed study of every constituent that enters into the value 57280 cal/gfw. For those constituents that rest upon a direct determination in terms of the 15°-calorie, the factor 4.185 is to be preferred; whereas a determination of the proper factor, or factors, to be used for those constituents that rest upon electrical measurements requires a knowledge both of the exact value of each of the units actually employed in the measurements, and of the numerical factor by means of which the data were converted to calories. Each of these may differ from observer to observer. Such detailed study has not been attempted by the present compiler.

M. Randall 50 accepts formulas (3) as satisfactory representations of the molecular specific heats, the unit being 4.182 joule/gfw H₂O.

$$(C_p)_{H_2} = 6.5 + 0.0009T \tag{3a}$$

$$(C_p)_{0p} = 6.5 + 0.0010T \tag{3b}$$

$$(C_p)_{H_2O} = 8.81 - 0.0019T + 0.00000222T^2$$
 (3c)

Whence,

$$\Delta C \equiv (C_p)_{\text{Ha}0} - [(C_p)_{\text{Ha}2} + 0.5(C_p)_{\text{Oa}}] = -0.94 - 0.0033T + 0.00000222T^2$$

$$= -3.931 - 0.0138T + 0.00000928_tT^2 \text{ joule/gfw-Ha}0$$

Accepting this as a satisfactory expression of ΔC when P is the pressure of water-vapor saturated at 25 °C, formula (4) is obtained:

$$(Q_T)_g = (Q_{208.1})_g - 1.703 + 0.393108(T/100) + 0.069003(T/100)^2 - 0.00309468(T/100)^3 \text{kj/gfw}$$
 (4)

⁴⁷ Randall, M., Int. Crit. Tables, 7, 226 (1930).

Reynolds, O., and Moorby, W. H., Phil. Trans. (A), 190, 301-422 (1897).
 Barnes, H. T., Idem, 199, 149-263 (1902); Proc. Roy. Soc. (London) (A), 82, 390-395 (1909).

⁵⁰ Randall, M., Int. Crit. Tables, 7, 231 (1930).

Taking $(Q_{298.1})_g = 241.930 \text{ kj/gfw}$, this becomes

$$(Q_T)_g = 240.227 + 0.393108(T/100) + 0.069003(T/100)^2 - 0.00309468(T/100)^2 \,\mathrm{kj/gfw}$$
 (5)

$$= 241.75 + 0.700758(t/100) + 0.0436485(t/100)^{2} - 0.00309468(t/100)^{8} \text{ kj/gfw}$$
(6)

the temperature being t °C = (273.1 + t) °K = T °K. These apply directly to a pressure of 23.76 mm-Hg, but insofar as all three gases may be regarded as ideal they apply to any pressure at which water-vapor can exist at t °C and the formulas for the specific heats remain valid.

Likewise, accepting Rossini's value for the heat of formation of water at 25 °C, but using formulas (7) for the specific heats and a lower value

Table 2.—Heat of Formation of Water-vapor

 $(Q_T)_y = (Q_{TP})_y$ = heat evolved in the formation of 1 gfw-H₂O of vapor at temperature T °K and pressure P from H₂ and O₂ at the same total pressure (P) and temperature; $(q_T)_g = (Q_T)_g/18.0154$ = heat of formation of 1 g H₂O; $(Q_T)_g = (Q_{273,1})_y + \Delta Q$.

With the exception of the last column, the tabular values are based on $(Q_{298.1})_g = 241.93 \text{ kj/gfw}$ and the values of the specific heat accepted by Randall (see text); the values in the last column have been computed by equation (8), (Chipman).

The values apply only to those pressures at which water-vapor can exist at the temperature considered, and at which the formula for ΔC is valid.

Unit of Q and $\Delta Q = 1$ kj/gfw-H₂O; of q = 1 kj/g. Temp. = t °C.

			Randall		Chipman
t	T	$(Q_T)_g$	$(q_T)_g$	ΔQ	$(Q_T)_{\varrho}$
0	273.1	241.75	13.419	0.00	241.96
15	288.1	241.86°	13.425	0.11	242.11
18	291.1	241.88	13.426	0.13	242.13
20	293.1	241.89	13.427	0.14	242.16
25	298.1	241.93b	13.429	0.18	242.20
50	323.1	242.11	13,439	0.36	242.44
100	373.1	242.49	13.460	0.74	242.91
200	473.1	243.30	13.505	1.55	243.80
300	573.1	244.16	13.553	2.41	244.62
500	773.1	245.96	13.653	4.21	246.07
1000	1273.1	250.03	13.879	8.28	248.61
1500	1773.1	251.64	13.968	9.89	249.59
2000	2273.1	248.47	13.792	6.72	248.99
2500	2773.1	238.20	13.222	-3.55	(246.83)
3000	3273.1	218.50	12.129	-23.25	(243.09)

 $^{^{\}circ}$ A. D. Crow and W. E. Grimshaw 52 state that the value 58 kcal/gfw (= 242.7 kj/gfw) at 15 $^{\circ}$ C is accepted by the Research Department, Woolwich, England.

 $[^]b$ W. F. Giauque and M. F. Ashley 63 derive the value 57.823 kcal/gfw (= 241.99 kj/gfw) at 25° C.

⁵¹ Chipman, J., Ind. Eng. Chem., 24, 1013-1017 (1932).

⁶² Crow, A. D., and Grimshaw, W. E., Phil. Trans. (A), 230, 39-73 (1931).

⁵⁸ Giauque, W. F., and Ashley, M. F., Phys. Rev. (2), 43, 81-82 (L) (1933).

(43.70 kj/gfw) for the latent heat of vaporization, J. Chipman ⁵¹ obtained an expression equivalent to (8).

$$(C_p)_{H_2} = 6.70 + 0.0007T \text{ cal/gfw}$$
 (7a)

$$(C_p)_{0_2} = 6.50 + 0.0010T \text{ cal/gfw}$$
 (7b)

$$(C_p)_{H_{20}} = 7.20 + 0.0027T \text{ cal/gfw}$$
 (7c)

$$\Delta C = -2.75 + 0.0015T \text{ cal/gfw} = -11.509 + 0.00628T \text{ j/gfw}$$

$$(Q_T)_g = 239.05 + 1.151(T/100) - 0.03139(T/100)^2 \text{ kj/gfw}$$

$$= 241.959 + 0.9795(t/100) - 0.03139(t/100)^2 \text{ kj/gfw}$$
(8)

valid for the range $300 \ge T \ge 2300$ °K.

Values computed by each of these formulas (6, 8) are given in Table 2.

Water: Heat of Formation.

The heat evolved in the formation of one gfw $(18.0156 \text{ g})^*$ of (liquid) water at 25 °C and 1 atm (A) from H_2 and O_2 at the same temperature and total pressure has been found by F. D. Rossini ⁵⁴ to be

$$(Q_{298.1,A})_w = 285.775 \pm 0.040 \text{ (Int.) kj} = 285.890 \text{ kj} \dagger$$

= 15.8626 (Int.) kj/g = 15.8690 kj/g

This is believed to be the best value at present available.

To the same heat of formation, F. R. Bichowsky ⁵⁵ has assigned the value 286.2 kj/gfw at 18 °C, based upon the work of various experimenters ⁵⁶⁻⁶⁸; and M. Randall the values 285.5 kj/gfw at 25 °C, 285.8 at 18 °C, and 286.3 at 0 °C. The sources upon which Randall's values rest have already been given (p. 14).

So far as H₂ and O₂ may be considered ideal and both the compressibility and the variation of the internal energy of water may be ignored, this same value will hold good for any pressure at which water can exist as a

*This is the value used by Rossini; it is 0.0002 g greater than that given in the Critical Tables and generally used in this compilation.

†This is negligibly affected by an error in reduction, the corrected value being 285.782 (Int.) kj = 285.897 kj [Rossini, F. D., Bur. Stand. J. Res., 7, 329-330 (RP343) (1931)].

⁵⁴ Rossini, F. D., Bur. Stand. J. Res., 6, 1-35 (RP259) (1931) → Proc. Nat. Acad. Sci., 16, 694-699 (1930) → Science, (N.S.) 72, 378 (1930).

55 Bichowsky, F. R., Int. Crit. Tables, 5, 176 (1929).

- 56 Abria, Compt. rend., 22, 372-373 (1846).
- ⁸⁷ Andrews, T., Phil. Mag. (3), 32, .321-339 (1848).
- 58 Berthelot, M., Ann. de Chim. et phys. (5), 23, 176-187 (1881).
- 59 Berthelot, M., and Matignon, C., Idem (6), 30, 547-565 (1893).
- ⁶⁰ Despretz, C., Idem (2), 37, 180-181 (1828).
- 61 Dulong, Compt. rend., 7, 871-877 (1838).
- 62 Favre, P. A., and Silbermann, J. T., Ann. de chim. et phys. (3), 34, 357-450 (1852).
- en Grassi, C., J. pharm. et chim. (3), 8, 170-181 (1845).
- 64 Mixter, W. G., Amer. J. Sci. (4), 16, 214-228 (1903).
- ⁶⁶ Rümelin, G., Z. physik. Chcm., 58, 449-466 (1907).
- 60 Schuller, A., and Wartha, V., Ann. d. Physik (Wicd.), 2, 359-383 (1877).
- or von Than, C., Ber. deut. chem. Ges., 10, 947-952 (1877); Ann. d. Physik (Wied.), 13, 84-105 (1881).
- **Thomsen, J., "Systematisk gennemförte termokemiske undersögelsers numeriske og teortiske resultater." 1882-1886.

liquid at 25 °C, and the value at any other temperature and pressure may be

determined by means of the relation
$$(Q_{TP})_w = (Q_{298.1,A})_w - \int_{298.1}^T (\Delta C)_w dT$$
,

the pressure being not less than that of water-vapor saturated at $T \circ K$.

No entirely satisfactory formulation for the specific heat of water at constant pressure is available, but for temperatures between 0 and 100 °C its variation is so slight (maximum 18.03, minimum 18.01 cal/gfw·°C) that the error introduced by regarding it as constant and equal to 18.02 may, for our present purposes, be ignored. By combining this expression with those for the specific heat of the gases (3) one obtains the expression $(\Delta C)_w = 8.27 - 0.0014T$ cal/gfw·°K = 34.58 - 0.00585T j/gfw·°K, the calorie being here taken as equivalent to 4.182 j (p. 15). In his compilation M. Randall ⁶⁹ has taken $(\Delta C)_w = 8.0$ cal/gfw·°K = 33.45 j/gfw·°K, independent of T. The first is to be preferred; it leads to the relation $(Q_{TP})_w = Q_{298.1,1})_w + 0.823 - 3.298(t/100) + 0.0292(t/100)^2$ kj/gfw, which becomes $(Q_{TP})_w = 286.713 - 3.298(t/100) + 0.0292(t/100)^2$ kj/gfw-H₂O when $(Q_{298.1,4})_w = 285.890$. Values computed by means of that formula are given in Table 3. Under the same conditions, Randall's expression for $(\Delta C)_w$ leads to $(Q_{TP})_w = 286.726 - 3.346(t/100)$ kj/gfw.

Table 3.-Heat of Formation of Water

 $(Q_{TP})_w$ = heat evolved in the formation of 1 gfw-H₂O of liquid at the temperature t and pressure P from H₂ and O₂ at the same temperature t and total pressure P. The pressure P may have any value at which water can exist as a liquid at temperature t.

These values have been computed by means of the more exact formula involving the square of t; by adding to each the associated δ , the corresponding value defined by the formula based on Randall's approximate value for $(\Delta C)_w$ will be obtained.

	Unit of (QTP)	w and of $\delta =$	1 kj/gfw-H ₂ O.	Temp. = t °C.	
*	$(Q_{TP})_{w}$	δ	ŧ	$(Q_{TP})_w$	8
0	286.713	0.013	30	285.726	-0.003
10	286.383	0.009	40	285.398	-0.010
15	286.219ª	0.006	50	285.071	-0.018
18	286.120	0.004	60	284.744	-0.026
20	286.054	0.003	80	284.093	-0.043
25	285.890	0	100	283.444	-0.064

^e A. D. Crow and W. E. Grimshaw ⁸² have stated that the value at 15 °C accepted by the Research Department, Woolwich, England, is 68.4 kcal/gfw (= 286.25 kj/gfw).

Ice: Heat of Formation.

From Rossini's value for the heat of formation of water (p. 17), the excess (0.82 kj/gfw, Table 3) of the heat of formation of water at 0 °C

 $^{^{\}infty}$ Randall, M., Int. Crit. Tables, 7, 232 (1930), the coefficient of $T \log T$ in the expression in the second line under $H_2O(l)$ at top of the page.

over that at 25 °C, and the latent heat of fusion of ice $(0.3336 \times 18.0154 = 6.01 \text{ kj/gfw}$, Table 272), it follows that the heat evolved in the formation of one gfw-H₂O of ice at 0 °C and one atmosphere, from gaseous H₂ and O₂ at the same temperature and pressure, is

$$(Q_{273.1,A})_i = 292.72 \text{ kj/gfw-H}_2\text{O}$$

= 16.248 kj/g

All of this refers to ice-I, the usual type that melts at 0 °C under a pressure of one atmosphere.

Insofar as H_2 and O_2 may be considered ideal, and both the compressibility and the variation of the internal energy of ice-I may be ignored, this same value will hold for any pressure under which ice-I can exist at 0 °C; and the value at any other temperature and pressure may be determined

from the relation $(Q_{TP})_i = (Q_{273.1,A})_i - \int_{273.1}^T (\Delta C)_i dT$, but only within the

domain in which ice-I can exist and within which the limitations just imposed are fulfilled.

Combining the expression $(0.5057 + 0.001863t \text{ cal}_{20}/\text{g} \cdot ^{\circ}\text{C})$ found by II. C. Dickinson and N. S. Osborne ⁷⁰ for the specific heat of ice with those (p. 15) for the specific heats of H₂ and O₂, one obtains $(\Delta C)_i = -4.282 + 0.1320t \text{ j/gfw} \cdot ^{\circ}\text{C} = -0.040331 + 0.01320(T/100) \text{ kj/gfw} \cdot ^{\circ}\text{K}$. Whence, $(Q_{TP})_i = (Q_{273 1,A})_i + 0.4282(t/100) - 0.660(t/100)^2 \text{ kj-gfw}$, which yields the following tabulated values when $(Q_{273,1,A})_i = 292.72 \text{ kj/gfw} \cdot \text{H}_2\text{O}$. In the range 0 to $-40 \, ^{\circ}\text{C}$ those values are only 0.01 j/gfw smaller than the corresponding ones based on the value of $(\Delta C)_i$ given in M. Randall's compilation; T^1 viz.,

$$(\Delta C)_i = (\Delta C)_w + (\Delta C)_{w \to i} = (+8.0 - 9.11 + 0.0336t)4.182$$

= $-4.642 + 0.1405t$ j/gfw.°C.

t 0 -10 -20 -30 -40 °C
$$(Q_{TP})_4$$
 292.72 292.67 292.61 292.53 292.44 kj/gfw-H₂O

6. Disposable Energy of Formation of H₂O at Constant Pressure

Of the greatest amount $(W_{TP} + P\Delta v)$ of external work that can be obtained from a process in which the final total pressure and temperature are the same as the initial ones, and the final volume exceeds the initial by the amount Δv , the amount W_{TP} is related to the corresponding heat evolution (Q_{TP}) of the process as indicated by the relation $(d/dT)(W_{TP}/T) = -Q_{TP}/T^2$. But of the total energy available under the best conditions within the stated limitations the amount $P\Delta v$ has to be expended in over-

⁷⁰ Dickinson, H. C., and Osborne, N. S., Bull. Bur. Stand., 12, 49-81 (SP248) (1915).

⁷¹ Randall, M., Int. Crit. Tables, 7, 232, 1930, $H_2O(l)$ and $H_2O(l) = H_2O(s)$.

coming the pressure, thus leaving only W_{TP} disposable for other work. For this reason, W_{TP} is sometimes called the technical work, or the technical energy, freed by the process; here it will be called the disposable energy. It is the decrement of the quantity (E + Pv - TS) that was denoted by ζ in Gibbs' papers, and that has been called the thermodynamic potential at constant pressure, the free energy at constant pressure, and the Gibbs function, but is called by G. N. Lewis and M. Randall ⁴⁶ simply the free energy, which term had already been used by Helmholtz and others to denote the quantity (E - TS). Here, E denotes the internal energy, E the entropy, and E0, E1, and E2 denote the pressure, volume, and absolute temperature, respectively.

It may readily be shown that
$$W_{TP} = W_{TP_1} - \int_{P_1}^{P} (\Delta v)_T dp$$
. If each

of the substances involved in the process is either an ideal gas or one having a volume that is negligible as compared with Δv , then $\Delta v =$ $(\Delta n)RT/P$, where Δn is the increase in the number of molecules of the gaseous substances per gfw of the compound formed, and $W_{TP} = W_{TP_1}$ -- $(\Delta n)RT\log_e(P/P_1)$. It is convenient, and a common practice, to split the logarithm into two parts, $\log(P/P_1) = \log(A/P_1) + \log(P/A)$, A denoting the pressure of one normal atmosphere, and to tabulate $w \equiv W_{TP_1} - (\Delta n)RT \log_t(A/P_1)$ instead of W_{TP_1} , the latter being given by the relation $W_{TP} = w - (\Delta n)RT \log_e(P/A)$. If the assumed phases with which we are concerned can exist under the pressure A when at the temperature T, then w is the value of W_{TP} at the pressure of one normal atmosphere; otherwise, as in the case of water-vapor at temperatures below 100 °C, w is a purely fictitious quantity, i.e., it is merely the numerical value obtained by adding to W_{TP} the numerical value of the quantity $(\Delta n)RT \log_e(P/A)$. The somewhat common practice of stating without reservation that w is the value of W_{TP} at one atmosphere is undesirable.

If $Q_{T_1P_1}$ and $W_{T_1P_1}$ are, respectively, the heat of formation and the disposable energy corresponding to T_1 and P_1 , and if $\Delta C = a + b_1T + b_2T^2 + \ldots b_nT^n$, then the value of W at the same pressure (P_1) and any temperature (T) within the allowable range is given by expression (9).

$$W_{TP_1} = Q_{T_1P_1} - (Q_{T_1P_1} - lV_{T_1P_1}) (T/T_1) + K - BT + f(T) + a\{T \log_{\bullet}(T/T_1) - (T - T_1)\}$$
(9)

where

$$K = \frac{1}{2}b_{1}T_{1}^{2} + \frac{1}{3}b_{2}T_{1}^{3} + \frac{1}{4}b_{3}T_{1}^{4} + \dots + \frac{1}{n+1}b_{n}T_{1}^{n+1}$$

$$B = b_{1}T_{1} + \frac{1}{2}b_{2}T_{1}^{2} + \frac{1}{3}b_{1}T_{1}^{3} + \dots + \frac{1}{n}b_{n}T_{1}^{n}$$

$$f(T) = \frac{1}{2}b_{1}T^{2} + \frac{1}{6}b_{2}T^{3} + \frac{1}{12}b_{3}T^{4} + \dots + \frac{1}{n(n+1)}b_{n}T^{n+1}$$

If $(T - T_1)$ is small, then $\{T \log_e(T/T_1) - (T - T_1)\}$ is small also, being equal to

 $T_1\left\{\left(1+\frac{T-T_1}{T_1}\right)\log_e\left(1+\frac{T-T_1}{T_1}\right)-\frac{T-T_1}{T_1}\right\}.$

Whenever $\Delta v = (\Delta n)RT/P$, the value of W for any associated temperture (T) and pressure (P) within the allowable range is given by the expression $W_{TP} = W_{TP_1} - (\Delta n)RT \log_e(A/P_1) - (\Delta n)RT \log_e(P/A)$, which is equivalent to (10), a form more convenient for computation.

$$W_{TP} = W_{T_{1}P_{1}} - (\Delta n)RT_{0}\log_{\bullet}(A/P_{1}) + aT_{0}\log_{\bullet}(T_{0}/T_{1}) + \tau\{a + (Q\tau_{1}P_{1} - W\tau_{1}P_{1})/T_{1}\} + \tau^{2}(\frac{1}{2}D_{1} + \frac{1}{3}D_{2}\tau + \dots + \frac{1}{n+1}D_{n}\tau^{n-1}) - t\left\{ (Q\tau_{1}P_{1} - W\tau_{1}P_{1})/T_{1} + (\Delta n)R\log_{\bullet}(A/P_{1}) - a\log_{\bullet}(T_{0}/T_{1}) + \tau(D_{1} + \frac{1}{2}D_{2}\tau + \dots + \frac{1}{n}D_{n}\tau^{n-1})\right\} + aT_{0}\left(\frac{T}{T_{0}}\log_{\bullet}\frac{T}{T_{0}} - \frac{t}{T_{0}}\right) + \frac{1}{2}D_{1}t^{2} + \frac{1}{6}D_{2}t^{2} + \dots + \frac{1}{n(n+1)}D_{n}t^{n+1} - (\Delta n)RT\log_{\bullet}(P/A)$$

$$(10)$$

where

If two phases, ϕ_1 and ϕ_2 , of a substance are in equilibrium at T' and P', then $(W_{T'P'})_{\phi_1} = (W_{T'P'})_{\phi_2}$ and $(Q_{T'P'})_{\phi_1} = (Q_{T'P'})_{\phi_2} + L_{12}$, where L_{12} is the latent heat absorbed by the substance in passing from phase ϕ_1 to phase ϕ_2 .

Quite recently, H. Zeise 72 has reviewed the several methods for obtaining w, and has derived the essential formulas.

Wherever it is necessary in the following paragraphs to know that the given data are expressed in terms of the international joule, as distinguished from the absolute, the symbol "Int." will accompany the symbol for the unit (see p. 13).

⁷² Zeise, H., Z. Elcktroch., 39, 758-773, 895-909 (1933).

Water-vapor: Disposable Energy of Formation.

In order to obtain from expressions (9) and (10) the value of W_{TP} , it is necessary to know the value of W_{TP} for some one set of associated values of P and T. Assuming the value. ($W_{298.1,A}$)_w = $56.560 \times 4.182 = 236.53 \text{ kj/gfw-II}_2\text{O}$, given by M. Randall ⁷¹ for the disposable energy in the formation of water at 25 °C and 1 atm, and remembering that at the pressure (23.76 mm-Hg) of water-vapor saturated at 25 °C the disposable energy of formation at 25 °C is the same for the vapor as for the liquid, one obtains the value ($W_{298.1,23.76\text{mm}}$)_g = 223.65 kj/gfw. Whence, taking

Table 4.—Disposable Energy in the Formation of Water-vapor

The greatest amount of external work that can be obtained from the combination of H_2 and O_2 at temperature T and total pressure P, to form water-vapor at the same temperature and pressure is $(H^r_{TP})_g + P\Delta v$, where Δv is the accompanying increase in volume, and P does not exceed either that at which water-vapor can exist at temperature T or that at which the formula used for ΔC is valid; $(H^r_{TP})_g \equiv v + 0.5RT \log_c(P/A)$, where w takes the values tabulated below, and A = pressure of one normal atmosphere; $w_g \equiv w_{12}/18.0154$; $P\Delta v = -0.5RT$.

Computations by means of formulas (12), (13), and (14) are indicated by subscripts; w_{12} is essentially the negative of M. Randall's ⁷¹ ΔF° . For values computed from molecular and spectroscopic data, see A. R. Gordon.⁸³

	Unit of w and	0.5RT - 1	kj/gfw-H2O;	of $w_{\theta} = 1 \text{ kj/g}$.	Temp.	- t ' C.
t	T	w_{12}	0.5RT	w _a	7013	v_{14}
0	273.1	229.11	1.135	12.718	229.25	229.12
15	288.1	228.41	1.198	12.679	228.55	228.41
18	291.1	228.27	1.210	12.671	228.40	228.27
20	293.1	228.18	1.218	12.666	228.30	228.18
25	298.1	227.95°	1.239	12.653	228.07	227.94
50	323.1	226.76	1.343	12.587	226.87	226.75
100	373.1	224.36	1.551	12.454	224.44	224.32
200	473.1	219.40	1.967	12.178	219.38	219.29
300	573.1	214.27	2.383	11.894	214.14	214.19
500	773.1	203.6	3.214	11.30	203.3	203.3
1000	1273.1	175.0	5.293	9.71	174.8	174.9
1500	1773.1	145.1	7.372	8.05	145.6	145.8
2000	2273.1	115.4	9.450	6.41	116.3	117.0
2500	2773.1	87.0	11.529	4.83	87.4	89.3
3000	3273.1	61.3	13.608	3.40	58.9	63.4

[&]quot;Randall gives 54.507 × 4.182 = 227.95 kj/gfw; W. F. Giauque and M. F. Ashley is give 54.670 × 4.185 = 228.79 kj/gfw at 25 °C; F. D. Rossini derives from his own value for the heat of formation of water (p. 17), together with various published spectroscopic values for the entropies of the gases, 228.57 (Int.) kj/gfw at 25 °C (=228.67 kj/gfw).

⁷³ Bjerrum, N., Z. physik. Chem., 79, 513-536 (1912).

⁷⁴ Langmuir, I., J. Am. Chem. Soc., 28, 1357-1379 (1906).

⁷⁵ Löwenstein, L., Z. physik. Chem., 54, 715-726 (1905).

⁷⁰ Nernst, W., and v. Wartenberg, H., Nachr. k. Ges. Wiss., Göttingen, 1905, 35-45 (1905).

⁷⁷ v. Wartenberg, H., Ber. deut. physik. Ges., 8, 97-103 (1906); Z. physik. Chem., 56, 513-533 (1906).

 $(Q_{298.1,23.76\text{mm}})_g = 241.93 \text{ kj/gfw}$ (p. 14), accepting Randall's expression for ΔC (p. 15), and using expression (9), one obtains

$$(W_{TP})_g = 240.227 - 0.003931T \log_e T - 0.0690(T/100)^2 + 0.0015473(T/100)^3 - 0.01688_5T + 0.004157_5T \log_e (P/A) \text{ kj/gfw-H}_2O$$
(11)

which differs from the one given by Randall solely by the presence of the last term and by small changes in the first and in the next to the last terms. The last two result from the use of a slightly different value for the heat of formation of water, and the first is necessary for the completion of the expression. Obviously, expression (11) ceases to have a physical significance when P exceeds the pressure of water-vapor saturated at T °K. The alternative form corresponding to (10) is

$$(W_{TP})_{g} = 229.110 - 0.04629_{1}t - 0.0562_{2} \left(\frac{t}{100}\right)^{2} + 0.001547_{3} \left(\frac{t}{100}\right)^{3} - 1.0736 \left\{\frac{T}{T_{0}} \log_{e} \left(\frac{T}{T_{0}}\right) - \frac{t}{T_{0}}\right\} + 0.004157_{5}T \log_{e} (P/A) \text{ kj/gfw-H}_{2}O \quad (12)$$

It seems that the expression given by Randall rests upon the value he gives for the heat of formation of water (p. 14), together with his conclusions regarding the thermal dissociation of water-vapor, as derived from the work of N. Bjerrum,⁷³ I. Langmuir,⁷⁴ L. Löwenstein,⁷⁵ W. Nernst and H. v. Wartenberg,⁷⁶ H. v. Wartenberg,⁷⁷ with a consideration of that of F. Haber and F. Fleischmann,⁷⁸ F. Faber and G. W. A. Foster,⁷⁹ A. Holt,⁸⁰ W. Nernst,⁸¹ and Schmidt,⁸² these being the references given in that section of his table.

Using other data (see p. 17), Chipman obtained an expression equivalent to (13)

$$(W_{TP})_g = 239.05 - 1.151(T/100) \log_c T + 0.03139(T/100)^2 + 2.783(T/100) + 0.004157_5 T \log_c (P/A) \text{ kj/gfw-H}_2\text{O}.$$
 (13)

=
$$229.25 - 4.6529(t/100) - 3.1434\{(1 + t/273.1) \log_e(1 + t/273.1) - (t/273.1)\} + 0.03139(t/100)^2 + 0.004157_5T \log_e(P/A) \text{ kj/gfw-H}_2O$$
,

and E. D. Eastman 82n one equivalent to (14)

$$(W_{TP})_g = 239.50 - 0.8412(T/100) \log_e T - 0.0090605(T/100)^2 + 0.0007366(T/100)^3 + 0.9374(T/100) + 0.004157_5 T \log_e (P/A) =$$

⁷⁸ Haber, F., and Fleischmann, F., Z. anorg. allgem. Chem., 51, 245-288 (1906).

⁷⁹ Haber, F., and Foster, G. W. A., Idem, 51, 289-314 (1906).

⁸⁰ Holt, A., Phil. Mag. (6), 13, 630-635 (1907).

⁸¹ Nernst, W., Z. anorg. allgem. Chem., 45, 126-131 (1905).

⁸² Schmidt, Diss., Berlin, 1921.

⁸²a Eastman, E. D., Bur. Mines, Circ. 6125: p. 15 (1929).

⁸⁸ Gordon, A. R., J. Chem'l Phys., 1, 308-312 (1933).

⁸⁴ Rossini, F. D., private communication, 1935.

$$229.12 - 4.6558(t/100) - 2.2973 \{(1 + t/273.1) \log_{e}(1 + t/273.1) - t/273.1\} - 0.003026(t/100)^{2} + 0.0007366(t/100)^{3} + 0.004157_{5}T \log_{e}(P/A) \text{ kj/gfw-H}_{2}O.$$
(14)

Values computed by each of these formulas are given in Table 4.

Water: Disposable Energy of Formation.

Accepting for the disposable energy in the formation of water at 25° C and 1 atm the value given by M. Randall, 71 viz., $(W_{298.1,A})_w = 56.560 \times 4.182 = 236.53_4$ kj/gfw-H₂O, for the heat of formation $(Q_{298.1,A})_w = 285.890$ kj/gfw-H₂O (p. 17), and for $(\Delta C)_w$ the expression $(\Delta C)_w = 34.58 - 0.00585T$ j/gfw-H₂O·°C (p. 18), one obtains expression (15), in which A = pressure of one normal atmosphere. It has no physical significance if P is less than the vapor-pressure of water at t°C.

$$(W_{TP})_w = 240.70_9 - 0.16845t - 0.00000292t^2 + 9.444 \{(1 + t/273.1) \times \log_e(1 + t/273.1) - t/273.1\} + 0.01247T \log_e(P/A) \text{ kj/gfw-H}_2O. (15)$$
 Values computed by means of (15) are given in Table 5.

Table 5.—Disposable Energy in the Formation of Water

The greatest amount of external work that can be obtained from the combination of H_2 and O_2 at temperature T and pressure P, not less than the vapor-pressure of water at $T \circ K$, to form water at the same temperature and pressure is $(W_{TP})_w + P(\Delta v) \equiv w + 1.5RT \log_e(P/A) + P(\Delta v)$, where A = pressure of 1 atm, and (Δv) is the increase in volume on passing from the mixed gases to the resultant water; $P(\Delta v) = -1.5RT = -0.01247T$ kj/gfw-H₂O. The following values of w have been computed by formula (15); $w_g = w/18.0154$.

nıt	or w a	nd 1.5K1 - 1	KJ/glw-112O, of w	, - I kj/g; temj	, , C - 1 I
	ŧ	au	w	1.5 <i>RT</i>	w_{g}
	0	273.1	240.709	3.406	13.3613
	10	283.1	239.031	3.531	13.2682
	15	288.1	238.196	3.593	13.2218
	20	293.1	237.363	3.656	13.1756
	25	298.1	236.535"	3.718	13.1296
	30	303.1	235.708	3.780	13.0837
	40	313.1	234.059	3.905	12.9922
	60	333.1	230.804	4.154	12.8115
	80	353.1	227.585	4.404	12.6328
	100	373.1	224.402	4,653	12.4561

Unit of w and $1.5RT = 1 \text{ kj/gfw-H}_2\text{O}$; of $w_g = 1 \text{ kj/g}$; temp. = $t \, ^{\circ}\text{C} = T \, ^{\circ}\text{K}$.

 $^{\circ}$ W. F. Giauque and M. F. Ashley 58 have concluded that $W=56.720\times4.185=237.37$ kj/gfw at 25 $^{\circ}$ C. Had this value been used instead of Randall's, the computed values of W would have exceeded those in the table by 0.77+0.0028t kj/gfw.

If for $(\Delta C)_w$ one uses Randall's approximation (33.45 j/gfw.°C, p. 18), keeping all else as before, one obtains a formula differing slightly

from (15), but yielding essentially the same values except at the higher temperatures. But even at 100 °C the value so obtained is only 0.006 kj/gfw smaller than that given in Table 5. It will be noticed that this difference at 100° C is only a tenth of that similarly produced in the computed value of the heat of formation (Table 3).

Ice: Disposable Energy of Formation.

We have found for water that $(W_{273.1,A}) = 240.71$ kj/gfw (Table 5); hence $(W_{273.1,A})_4 = 240.71$ kj/gfw also, water and ice-I (the common type of ice) being in equilibrium at that temperature and pressure. The heat of formation of ice is $(Q_{273.1,A})_4 = 292.72$ kj/gfw (p. 19), and $(\Delta C)_4 = -0.040331 + 0.01320(T/100)$ kj/gfw °K (p 19). Hence the disposable energy in the formation of ice from H_2 and O_2 at the same T and P may be computed by means of (16).

$$(W_{TP})_{i} = 240.71 - 0.1904t - 11.014 \{ (T/T_{0}) \log_{e}(T/T_{0}) - t/T_{0} \} + 0.660(t/100)^{2} + 0.01247T \log_{e}(P/A)$$

$$\equiv w + 0.01247T \log_{e}(P/A) \text{ kj/gfw-H}_{2}O \qquad (16)$$

 $T_0 = 273.1$, T = 273.1 + t, temperature = t °C, A = pressure of one normal atmosphere. This expression has no physical significance at temperatures above the melting point of ice under pressure P, but may be extrapolated to w = 235.94 for T = 298.1, which is the same as the value $(56.418 \times 4.182 = 235.94)$ given in M. Randall's compilation.

IB. DISSOCIATION

7. DISSOCIATION OF WATER-VAPOR

Thermal Dissociation of Water-vapor.

When a mixture of H_2 , O_2 , and H_2O -vapor is in thermal equilibrium, $k_1(n_w/v)^2 = k_2(n_H/v)^2(n_O/v)$, where v is the volume of the mixture; n_w , n_H , and n_O are, respectively, the number of moles of H_2O , H_2 , and O_2 ; k_1 is half the number of moles of H_2O -vapor that dissociate in unit time when the concentration (n_w/v) is unity; and k_2 is half the number of moles of H_2O formed per unit of time in a mixture of H_2 and H_2 and H_2 in which the concentration $(n_H/v, n_O/v)$ of each gas is unity. The factor $\frac{1}{2}$ arises from the fact that two molecules of H_2O are necessarily involved in each case, the reaction being $2H_2O \rightleftharpoons 2H_2 + O_2$. The values of the k's vary with the temperature and with the unit of concentration, but, to at least a first approximation, are independent of the actual concentrations. The numerical value of k_1 varies as the square of the size of the unit of concentration, and that of k_2 as the cube.

Their ratio $(K = k_1/k_2)$ is the ratio of the number of moles of H_2O -vapor that dissociate to the number formed in the same time when the concentration of each of the three gases is unity. It is called the *constant of dissociation*, and in this case it has the dimensions of a concentration. The reciprocal of K might logically be called the constant of combination, but unfortunately, it also is occasionally called the constant of dissociation. Obviously, any function of K and the temperature is an isothermal constant, but that does not justify the confusion introduced by calling such arbitrary functions constants of dissociation. Here we shall restrict that term to the ratio k_1/k_2 and shall denote it by K; $K = (n_H)^2(n_Q)/(n_w)^2v$.

Insofar as the three gases may be considered ideal, this expression for K is equivalent to (17) in which the p's indicate the partial pressures.

$$KRT = (p_H)^2 (p_0) / (p_w)^2$$
 (17)

The product KRT, which in this case is of the dimensions of a pressure, is frequently denoted by K_p and called the dissociation constant; and so is its square root. We shall not use that term to denote either KRT or $(KRT)^{\frac{1}{2}}$.

If the mixed gases have been derived from the dissociation of W moles of H_2O , and if α is the fraction of W dissociated when the mixture is in equilibrium, then $n_w = (1 - \alpha)W$, $n_H = \alpha W$, $n_O = \alpha W/2$, and the relations already found become

$$K = \frac{\alpha^3}{2(1-\alpha)^2} \left(\frac{W}{v}\right) \tag{18}$$

$$KRT = \frac{\alpha^3}{(1-\alpha)^2(2+\alpha)} (P)$$
 (19)

Here, P is the total pressure, and 18.0154W/v is the actual density of the mixture. The first of these expressions (18) is always true if the reaction is $2H_2O \rightleftharpoons 2H_2 + O_2$ and the relation stated at the beginning of this section applies; but the second (19) can apply only in the domain in which the gases may be regarded as ideal. The computation of α when the value of KRT/P is known may be facilitated by the use of Table 7.

If both sides of (18) are divided by C_1 (= the concentration of one gfw per liter) they become dimensionless; similarly if (19) is divided by A, the pressure of one normal atmosphere. It may be shown that $(d/dT)(\log_{\theta}KRT/A) = 2(Q_{TP})_{g}/RT^2 = -2(d/dT)(W_{TP}/RT)_{g}$, where $(Q_{TP})_{g}$ and $(W_{TP})_{g}$ have each the same significance as on pp. 12, 19. Whence from (11) one obtains (20) for P = A,

$$-\frac{RT}{2}\log_{\theta}\left(\frac{KRT}{A}\right) = 240.247 - 0.003931T\log_{\theta}T - 0.0690\left(\frac{T}{100}\right)^{2} + 0.001548_{3}\left(\frac{T}{100}\right)^{3} - I_{a}T \text{ kj/gfw-H}_{2}O$$
(20)

where I_d , a constant of integration, is the same quantity as that designated by I in M. Randall's compilation, but is expressed in a different unit.

By means of this formula and the values of KRT/A derived from the observed values of α , the corresponding values of I_a can be computed; $R=8.315 \text{ j/gfw} \cdot {}^{\circ}\text{K}=0.8206 \text{ l·atm/gfw} \cdot {}^{\circ}\text{K}$. These values of I_a are given in Table 6; their mean is 15.54 j/gfw $\cdot {}^{\circ}\text{K}$. By means of it and relation (20) values of K/C_1 and of α for P=A have been computed; they also are given in the **table**.

B. Lewis and J. B. Friauf 2 have derived the following expressions:

$$0.5 \log_{10}(A/KRT) = \frac{57295}{4.573T} - 0.848 \log_{10}T - 1.474 \frac{T}{10000} +$$

$$7.78 \left(\frac{T}{10000}\right)^{2} - 8.72 \left(\frac{T}{10000}\right)^{3} + 0.0616; T \ge 2800 \text{ °K}$$

$$0.5 \log_{10}(A/KRT) = \frac{14412}{T} + 2.0975 \log_{10}T - 0.7610 \frac{T}{10000} -$$

$$2.047 \left(\frac{T}{10000}\right)^{2} + 2.473 \left(\frac{T}{10000}\right)^{3} - 10.4417; 2800 \text{ °K} \ge T \ge 3000 \text{ °K}$$

$$0.5 \log_{10}(A/KRT) = \frac{13317.3}{T} - 0.34492 \log_{10}T - 4.3285;$$

$$T \ge 3000 \text{ °K}$$

$$(23)$$

Values computed by these equations are given in the second section of Table 6; for each of the temperatures 2800 and 3000, values have been computed by each of the equations applicable thereto.

The preceding discussion relates only to the thermal dissociation of water-vapor into uncharged molecules of H₂ and O₂. The questions arise: Does an increase in temperature give rise to other types of dissociation? Are any of the products of dissociation electrically charged?

- K. F. Bonhoeffer and H. Reichardt ⁸ have concluded, from their study of the thermal variation in the optical absorptivity of water-vapor, that at temperatures above 1200 °C the amount of vapor dissociating into $H_2 + 2OH$ is a little greater than that into $2H_2 + O_2$. The OH was electrically neutral.
- G. M. Woods and T. C. Poulter ⁴ found no detectable electrical conduction through water-vapor heated to 500 °C. The sensitivity of the method employed was not great.

¹ Randall, M., Int. Crit. Tables, 7, 231 (1930).

² Lewis, B., and Friauf, J. B., J. Am. Chem. Soc., 52, 3905-3920 (1930).

⁸ Bonhoeffer, K. F., and Reichardt, H., Z. physik. Chem. (A), 139, 75-97 (1928).

⁴ Woods, G. M., and Poulter, T. C., Proc. Iowa Acad. Sci., 33, 172-173 (1926).

Table 6.—Thermal Dissociation of Water-vapor

Values given in the first section of the main table are based upon the observational data used in M. Randall's compilation. Those data have been checked against the original publications for errors in transcription, and all computations have been made independently for this table. The quantity $(A/KRT)^{0.5}$ is denoted by K in Randall's compilation, where it is called the constant of dissociation.

 α = ratio of the number of gfw-H₂O dissociated into H₂ + 0.5O₂ to the total number of gfw of H₂O contained in a mixture of the three gases when the mixture is in equilibrium at the pressure and temperature indicated, and has the analytical composition H₂O. $K = (n_H)^2(n_0)/(n_w)^2v$, where v = total volume; n_w , n_H , and n_0 = the number of gfw of H₂O, H₂, and O₂, respectively, contained in that volume; A = the pressure of one normal atmosphere; C_1 = the concentration of one gfw per liter, the gases being assumed to be ideal; and I_d is a constant of integration (see text). The computed values (comp) have been derived from equation (20) with I_d = 15.54, its mean value. K/C_1 and KRT/A are each dimensionless; obviously, they are, respectively, the numerical values of K and of KRT when the unit of K is 1 gfw/l and that of KRT is 1 atm, but in writing such equations as (20) the A and the C_1 must be retained if mathematical nonsense is to be avoided.

D. M. Newitt ⁵ has given the following values for the high temperatures and pressures encountered in explosions of mixtures of H_2 and O_2 . P_i , P_m = initial and maximum pressures, respectively; T_m = maximum temperature; α_m = value of α corresponding to P_m and T_m . The values of K have been computed by the compiler from the tabulated values of P_m , T_m , and α_m .

P_{i}	3	10	25	50	75	100	125	150	175	atm
P_m	23	78	198	395	602	808	1010	1200	1400	atm
T_m	2585	2630	2660	2655	2700	2715	2720	2680	2690	°K
$100 \alpha_m$	2.2	1.8	1.7	1.4	1.4	1.5	1.3	1.2	1.0	
$10^{6}K/C_{1}$	0.6	1.1	2.3	2.6	5.2	6.3	5.1	4.8	3.4	

Newitt regards these values as superior to all earlier ones. In the computation of T_m and α_m , he introduced a correction for the variation of the co-volume with the temperature.

K. F. Bonhoeffer and H. Reichardt ³ have given the following values for the dissociations $H_2O \rightarrow 0.5H_2 + OH$ (subscript OH) and $H_2O \rightarrow H_2 + 0.5O_2$ (subscript O). The latter essentially agree with those of the main table.

T	1000	1300	1500	1705	1900	2155	2505	° K
$0.5 \log_{10}(A/KRT)_{OH}$	10.5	7.15	5.70	4.5_{0}	3.6_{0}	2.75	1.78	
$0.5 \log_{10}(A/KRT)_0$	10.0 ₅	7.00	5.71	4.64	3.80	3.04	2.15	

Table 6—(Continued)

For values computed from molecular and spectroscopic data, see A. R. Gordon.6

Unit of I = 1 j/gfw. K. Temp. = T K. A/KRT and K/C_1 are dimensionless.

				$= 10^n B$	1							
	1 atm	Va	lue	logio		K	$/C_1 = 1$	$0^{-n}B'$	$-\log_{10}($	K/C_1	1 atm	
T	100aobs	\boldsymbol{B}	92	0.0	I	n	B'obs	B'_{comp}	Obs	Comp	100 acomp	Ref.
1205	0.00325	7.63	6	6.883	14.85	16	1.58	1.86	15.801	15.731	0.00343	La
1325			6								0.00343	La
1354	0.0049	4.12		6.615	15.93	16	5.30	4.82	15.276	15.317		La
1393	0.0069	2.47	6	6.393	15.01	15	1.43	1.63	14.843	14.788	0.00719	
1397	0.0078	2.05	6	6.312	16.04	15	2.08	1.84	14.683	14.736	0.00750	йM
1433	0.0103	1.35	6	6.131	14.99	15	4.65	5.30	14.333	14.275	0.0108	La
1455	0.0142	8.36	5 5 5	5.922	16.35	15	12.0	9.88	13.921	14.005	0.0133	La
1474	0.0141	8.45	5	5.927	14.04	14	1.16	1.66	13.936	13.779	0.0159	La
1480	0.0189b	5.44	5	5.736	17.00	14	2.78	1.96	13.555	13.708	0.0168	NW
1531	0.0255	3.47	5	5.540	15.08	14	6.61	7.38	13.180	13.132	0.0265	La
1550	0.0287	2.91	5	5.464	14.53	14	9.29	11.82	13.032	12.927	0.0311	La
1561	0.034	2,26	5	5.354	15.50	13	1.54	1.54	12.814	12.811	0.0341	NW
1705	0.102	4.34	4	4.638	15.60	12	3.80	3,74	11.421	11.428	0.1014	Lo
1783	0.182	1.82	4	4.260	16.36	11	2.07	1.69	10.684	10,772	0.170	Lo
1863	0.354	6.71	3	3.827	18.60°	10	1.46	0.69	9.838	10.159	0.277	Lo
1968	0.518	3.78	3	3.578	16.18	10	4.33	3.71	9.363	9.430	0.492	Lo
2155	1.18	1.09	3	3.039	15.31	9	4.73	4.99	8.325	8.302	1.20	W
2257	1.77	5.93	2	2.773	15.31		1.54	1.62	7.814	7.790	1.80	
2337	2.8	2.95	2	2,470	17.35	8	5.98	3.84	7.222	7.412	2.43	S
2505	4.5	1.43	2 2 2	2.155	16.31	7	2.38	1.98	6,623	6.704	4.23	Š
2642	4.3	1.53	2	2.185	10.70	ź	1.97	6.30	6.705	6.201	6.28	Ř
2684	6.2	8.70	1	1.940	13.95	7	6.00	8.74	6.222	6.058	7.01	ć
2698	7.5	6.46	1	1.810	15.98	ź	10.8	9.72	5.966	6.012	7.27	n
2731	8.2	5.65	1	1.752	16.02	6	1.40	1.04	5.854	5.905	7.89	5
2761	6.6	7.96	1	1.901	12.210	6		1.55	6.157	5.811		D D
2834			1				0.70				8.49	WSSBSBSBBBB BBBBBBBBBBBBBBBBBBBBBBBBBBB
	9.8	4.26	1	1.629	15.21	6	2.37	2.57	5.625	5.590	10.05	P
2929	11.1	3.50	ī	1.544	14.15	6	3.40	4.75	5.469	5.323	12.3	B
3092	13.0	2.71	1	1.433	12.140	6	5.36	12.16	5.270	4.915	16.7	3
				Mean	15.54							

B. Lewis' and J. B. Friauf's 2 formulas (see p. 27) yield the following values.

			10 ⁿ B	K	$C_1 = 10$	nB'	
T	Valu	e-n	\log_{10}	B' Val	n	$-\log_{10}$	1 atm 100α
1000	9.30	9	9.968	1.41	21	20.851	2.85×10^{-5}
1200	6.48	7	7.811	2.42	18	17.616	7.81×10^{-4}
1400	1.842	6	6.2652	2.57	15	14.591	8.38×10^{-8}
1600	1.267	5	5.1028	4.74	13	12.324	0.0499
1800	1.576	4	4.1976	2.73	11	10.565	0.201
2000	2.974	3	3.4734	6.89	10	9.162	0.609
2200	7.62	2	2.8817	9.55	9	8.020	1.51
2400	2.451	2	2.3894	8.45	8	7.073	3.20
2500	1.490	2	2.1733	2.194	7	6.6587	4.39
2600	94.2	0	1.9739	5.29	7	6.2769	5.91
2800	41.6	0	1.6189	2.517	6	5.5991	9.95
2800	41.4	0	1.6165	2.545	6	5.5943	9.98
2900	28.7	0	1.4577	5.11	6	5.2919	12.55
3000	20.4	0	1.3098	9.75	6	5.0109	15.4
3000	20.4	0	1.3099	9.75	6	5.0111	15.4
3200	11.0	0	1.0421	3.14	5	4.5035	22.3
3500	5.00	0	0.6988	1.39	4	3.8558	34.4
4000	1.75	0	0.2432	9.94	4	3.0026	55.0

⁵ Newitt, D. M., Proc. Roy. Soc. (London) (A), 119, 464-480 (1928). ⁶ Gordon, A. R., J. Chem'l Phys., 1, 308-312 (1933).

Table 6 -- (Continued)

B. Lewis and G. von Elbe ⁷ have given the following values, with which those given by E. Justi and H. Lüder ⁸ essentially agree. Each denotes $(KRT/A)^{0.5}$ by the symbol K_p . Two types of dissociation are considered: $H_2O \rightleftharpoons H_2 + \frac{1}{2}O_2$ and $H_2O \rightleftharpoons \frac{1}{2}H_2 + OII$. The first is denoted in the table by the symbol H, O; the second by H, OH.

		(A/KRT)	$0.5 = 10^n B$			
	II, ()—	II, OI	I——	$\log_{10}(A/$	$KRT)^{0.5}$
T	\boldsymbol{B}	n	B	11	H, O	н, он
300	5.9	39	2	43	39.77	43.3
400	1.82	29	5	31	29.26	31.7
600	4.36	18	1	20	18.64	20.0
800	1.90	13	1.17	14	13.28	14.07
1000	1.12	10	3.39	10	10.05	10.53
1200	7.94	7	1.48	8	7.90	8.1 7
1400	2.19	6	2.95	6	6.34	6.47
1600	1.58	5	1.58	5	5.20	5.20
1800	1.86	4	1.55	4	4.27	4.19
2000	3.31	3	2.51	3	3.52	3.40
2200	8.13	2	5.50	2	2.91	2.74
2400	2.57	2	1.55	2	2.41	2.19
2600	1.00	2	5.50	1	2.00	1.74
2800	4.27	1	2.19	1	1.63	1.34
3000	2.04	1	9.98	0	1.31	0.999

a References:

Table 7.—Variation of α with KRT/P

From (19), $\alpha^3 = (1-\alpha)^2(2+\alpha)(KRT/P)$. If α is small, α^3 varies as 2KRT/P. If α exceeds 4 per cent, its approximate value can be read from this table, and the computation of a more exact value is facilitated by using in the calculation the approximate value here given for $(1-\alpha)^2(2+\alpha)$. For example, if KRT/P is 0.645, one sees at once that α lies between 62 and 63 per cent, and that $(1-\alpha)^2(2+\alpha)$ is about 0.374. Whence $\alpha^3 = 0.374(0.645)$ or $\alpha = 62.2$ per cent. Should greater precision be desired, use this value of α to compute $(1-\alpha)^2(2+\alpha)$, finding 0.3746; substitute this value in the cubic, and solve, finding $\alpha = 62.28$. Take the mean of this and the value (62.2) previously found, which gives $\alpha = 62.24$ per cent.

B = Bjerrum, N., Z. physik. Chem., 79, 513 536 (1912).

La = Langmuir, I., J. Am. Chem. Soc., 28, 1357-1379 (1906).

Lo = Löwenstein, L., Z. physik. Chem., 54, 715-726 (1905).

NW - Nernst, W., and Wartenberg, H. v., Nachr. v. d. k. Ges. Wiss. Göttingen, 1905, 35-45 (1905).

S = Siegel, W., Z. physik. Chem., 87, 641-668 (1914).

W = von Wartenberg, H., Ber. deut. physik. Ges., 8, 97-103 (1906).

b Frequently quoted incorrectly as 0.0184.

Omitted in taking the mean.

⁷ Lewis, B., and von Elbe, G., J. Am. Chem. Soc., 57, 612-614 (1935).

⁸ Justi, E., and Lüder, H., Forsch. Gebiete Ingenieurw., 6, 209-216 (1935).

7. DISSOCIATION OF VAPOR

Table 7—(Continued)

	-			-	
KRT/P	and	α	are	each	dimensionless.

	KKI/P and a	are each dimens	ioniess.	
KRT/P	P/KRT	$(P/KRT)^{0.8}$	$(1-\alpha)^2(2+\alpha)$	100α
34.05×10^{-6}	29370	171.4	1.880	4
1.186×10^{-4}	8430	91.8	1.820	4 6
2.909×10^{-4}	3440	58.6	1.760	8
4.210×10^{-4}	2375	48.7	1.731	ŏ
4.210 X 10			1.701	9 10
5.879×10^{-4} 7.96×10^{-4}	1701	41.2 35.44	1.671	11
7.90 X 10 °	1256	35.44	1.642	12
0.001051	951	30.84	1.612	12 13
0.001363	734 576	27.09	1.582	14
0.001735	5/0	24.00	1.553	15
0.002173	460	21.45	1.524	16
0.002688	372 301	19.29 17.35	1.481	17
0.003317		15.86	1.466	18
0.003975	251.6	14.48	1.437	10
0.004770	209.6		1.408	19 20
0.00568	176.0	13.27	1.379	21
0.00671	149.0	12.21 11.26	1.351	22
0.00788	126.9	10.42	1.322	23
0.00921	108.6	9.64	1.294	24
0.01076	92.9 81.0	9.04	1.266	25
0.01234		8.39	1.200	26
0.01420	70.4		1.238 1.210	26 27
0.01628	61.4	7.84 7.33	1.182	28
0.01857	53.8	7.33	1.154	20
0.02112	47.3 41.7	6.88	1.127	29 30
0.02396	41./	6.46	1.100	31
0.02708	36.93	6.077	1.073	32
0.03054	32.74 29.11	5.722 5.395	1.046	32 33
0.03435	29.11		1.019	34
0.03859	25.91	5.090	0.9929	35
0.04319	23.15	4.811	0.9666	36
0.04828	20.71	4.551	0.9406	37
0.0539	18.55	4.307	0.9149	38
0.0600	16.67	4.083		39
0.0668	14.98	3.870 3.674	0.8893 0.8640	40
0.0741	13.50	3.0/4	0.8389	41
0.0812	12.32 10.99	3.508 3.315	0.8141	42
0.0910	10.99	3.151	0.7895	43
0.1007	9.93	2.997	0.7652	44
0.1113	8.98	2.851	0.7411	45
0.1230	8.13	2.715	0.7173	46
0.1356	7. 37 6.68	2.584	0.6938	47
0.1496	6.06	2.462	0.6706	48
0.1650	5. 5 0	2.402	0.6476	49
0.1818	5.00	2.345 2.236	0.6250	50
0.2000	4.54	2.131	0.6026	51
0.2202	4.13	2.032	0.5806	52
0. 24 22 0.2 663	3.76	1.939	0.5589	52 53
	3.41	1.847	0.5575	54
0.2930	3.105	1.762	0.5575 0.5164	55
0.3220	2.821	1.680	0.4956	56
0.3545	2.567	1.602	0.4752	54 55 56 57 58
0.3896	2.30/	1.527	0.4752	£0
0.4287	2.333 2.119	1.456	0.4354	59
0.4720	2.119 1.926	1.388	0.4354	60
0.5192	1.926 1.748	1.322	0.3970	61
0.5720		1.260	0.3783	61 62
0.6300	1.587 1.439	1.200	0.3763	63
0.6948	1.439	1.142	0.3421	64
0.7662	1.303	1.142	0.3421	04

Table 7(Continued)										
KRT/P	P/KRT	$(P/KRT)^{0.5}$	$(1-\alpha)^2(2+\alpha)$	100α						
0.8460	1.182	1.087	0.3246	65						
0.9352	1.069	1.034	0.3075	66						
1.035	0.966	0.983	0.2908	67						
1.146	0.873	0.934	0.2744	68						
1.271	0.787	0.887	0.2585	69						
1.411	0.709	0.842	0.2430	70						
1.571	0.636	0.798	0.2279	71						
1.750	0.571	0.756	0.2132	72						
1.955	0.512	0.715	0.1990	73						
2.189	0.457	0.676	0.1852	74						
2.453	0.408	0.638	0.1719	75						
2.760	0 362	0 602	0.1590	76						
3.115	0.321	0.567	0.1465	77						
3 522	0.284	0.533	0.1346	78						
4.008	0.250	0.500	0.1230	79						
4.602	0.217	0.466	0.1120	80						
9.575	0 104	0.323	0.0641	85						

Table 8.—Thermal Dissociation of Water-vapor: Effect of Pressure and Temperature

The data in this table have been computed by means of equations (19) and (20) with $I_d = 15.54$, the mean of Table 6; they are valid only to the extent to which the constants in equation (20) are valid. The compiler expresses no opinion regarding their validity at the higher temperatures, but the values of α here given essentially agree with those given by N. Bjerrum in a similar table covering the same domain. See also Newitt's values at the head of Table 6. G. Chaudron 10 has expressed the opinion that the specific heats of the gases at high temperature are not known with sufficient precision to enable one to infer with certainty the dissociation at such temperatures. 11

P= total pressure; $\alpha=$ fraction of H_2O -vapor dissociated when the analytical composition of the mixed gases is that of H_2O ; K= constant of dissociation (see Table 6); $C_1=$ concentration of 1 gfw/l.

	Unit of P	= 1 atm. K/C_1	is dimensionless.	Temperature = T °K.	
	$P \rightarrow$	0.1	1	10	100
T	$-\log_{10}(K/C_1)$			100α	
1000	21.964	0.000056	0.000026	0.0000121	0.0000056
1500	13.478	0.0434	0.0202	0.00936	0.00434
2000	9.225	1.24	0.579	0.269	0.125
2500	6.724	8.77	4.17	1.96	0.914
3000	5.139	27.7	14.1	6.85	3.24
3500	4.113	51.1	29.4	15.1	7.34
4000	3.465	67.8	44.1	24.4	12.2
4500	3.098	76.6	54.7	31.7	16.5
5000	2 950	80.	58.7	35.6	18.8

Bjerrum, N., Z. physik. Chem., 79, 513-536 (1912).

¹⁰ Chaudron, G., Bull. soc. chim. de France (4), 37, 657-679 (1925).

¹¹ See also Gordon, A. R., and Barnes, C., J. Phys'l Chem., 36, 1143-1151 (1932).

Photochemical Dissociation of Water-vapor.

A. Coehn and G. Grote 12 have reported that under the radiation from a Hg-arc in a quartz tube the equilibrium amount of dissociation of water-vapor at a pressure of 0.825 atm is the same at 240 °C as at 150 °C, *i.e.*, 0.124 per cent, which is the same as for thermal dissociation at about 1730 °C. They give the following values for 150 °C, P being the initial pressure of the water-vapor, and α being the fraction dissociated when equilibrium is attained:

P	0.825	0.585	0.448	0.370	atm
α	0.1237	0.1992	0.2960	0.3618	per cent
$P\alpha$	0.102	0.116	0.132	0.135	-

From these data, which may be represented by the formula $P(\alpha + 0.0681) = 0.1600$, the observers infer that the reaction is of the first order. Two hours were required for establishing equilibrium.

R. S. Mulliken ¹³ has suggested that the strong continuous absorption band beginning suddenly near $\lambda = 1800 \text{A}$ and initiating the ultraviolet absorption of water-vapor represents a dissociation or a predissociation process.

H. Senftleben and I. Rehren ¹⁴ and J. R. Bates and H. S. Taylor ¹⁵ observed a dissociation into H and OH when a mixture of the vapors of water and of mercury was illuminated by the light from a suitable line of the mercury spectrum (cf. p. 60).

A. Tian ¹⁶ is occasionally quoted as having studied the photochemical dissociation of water-vapor, but the object of his investigations was liquid water, and he was of the opinion that the vapor was not involved (cf. p. 35).

A. Terenin and H. Neujmin ¹⁷ and H. Neujmin and A. Terenin ¹⁸ have reported that under the action of radiation in the range $\lambda=1300$ to 1500A, the excited OH radical is split off from the water-vapor molecule, carrying an abnormally great amount of rotational energy; and that the threshold energy for that dissociation is 207 kcal/gfw-H₂O (= 866 kj/gfw-H₂O).

Ionic Dissociation of Water-vapor. See also Sections 10 and 24.

In 1925, F. Hund ¹⁰ calculated the energy that would be required to remove one or both H-ions from H_2O constructed like his atomic model (p. 45), finding 370 ± 30 and 920 ± 40 kcal/gfw, respectively. These values are equivalent to 16.0 and 39.8 electron-volts per molecule or 1550 and 3850 kj/gfw. Recently, J. C. Slater ²⁰ has concluded that the energy of such a triangular molecule is -9.3 electron-volts.

¹² Coehn, A., and Grote, G., Nernst, "Festschrift," p. 136-167, 1912.

¹⁸ Mulliken, R. S., J. Chem'l Phys., 1, 492-503 (1933).

¹⁴ Senftleben, II., and Rehten, I., Z. Physik., 37, 529-538 (1926).

¹⁸ Bates, J. R., and Taylor, H. S., J. Am. Chem. Soc., 49, 2438-2456 (1927).

¹⁶ Tian, A., Compt. rend., 152, 1012-1014 (1911); Ann. de Phys. (9), 5, 248-365 (1916).

¹⁷ Terenin, A., and Neujmin, H., J. Chem'l Phys., 3, 436-437 (L) (1935).

¹⁸ Neujmin, H., and Terenin, A., Acta Physicochim. URSS, 5, 465-490 (1936).

K. F. Bonhoeffer 21 and K. F. Bonhoeffer and H. Reichardt 8 have obtained the following values: H₂ + 2OH = 2H₂O + 128 kcal; H + H = $H_2 + 101$ kcal. They concluded that the dissociation $H_2O \rightarrow H + OH$ requires 111 or 115 kcal/gfw = 464 or 481 kj/gfw = 4.8 or 5.0 electronvolts per formula molecule. But E. Gaviola and R. W. Wood 22 found that a greater value, probably about 5.2 electron-volts per molecule, is required for

Table 9.—Dissociation of Water-vapor in the Glow Discharge (See also Table 75)

A. Güntherschulze and H. Schnitger 29 give the following data for the dissociation 2H₂O

≥ 2H₂ + O₂ in the glow discharge, primarily in the cathode portion. Partial pressure of H_2O is p_1 , of the $(2H_2 + O_2)$ is p_2 ; T °K is the temperature at which the thermal dissociation at those pressures would equal that actually observed in the glow discharge at the temperature t °C; α is the dissociation coefficient as already defined (p. 26).

Unit of p_1 and of	$p_2 = 1$ mm-Hg; α	is dimensionless.	Temp. = t °C; for	T see heading
t	<i>p</i> ₁	p ₂	100α	T
-47.0	42	6.1	8.8	1915
-42.0	74	18.1	14.0	2095
-37.0	134	46.6	18.9	2170
-32.0	227	70.0	17.1	2190
-27.0	387	78.0	11.8	2120
-22.0	636	117.4	11.0	2140
-17.0	1027	145.	8.6	2115
-12.0	1627	329.	11.9	2245
0.0	4579	1078.	13.6	2380
+17.0	14530	5720.	24.9	275 0
18.2	15670	4850.	17.1	2600
22 .6	20560	10070.	24.8	2810

 $H_2O \rightarrow H + OH$, and that such dissociation occurs only once in about ten thousand collisions between the molecules of H₂O and of excited Hg. And more recently, H. Senftleben and O. Reichemeier ²³ have reported 5.05 ± 0.04 electron-volts for $H_2O \rightarrow H + OH$, and that the value increases with

R. S. Mulliken 24 has concluded from thermal data that the work per H-atom required to ionize H_2O completely is $110 \times 4.185 = 460$ ki, or 920 kj/gfw-H₂O. M. Magat ²⁵ from spectroscopic data derived for H₂O \rightarrow $2H^+ + O^-$ the energy $(269 \pm 3) \times 4.185 = 1126 \text{ kj/gfw}$, and from thermal data $218 \times 4.185 = 912 \text{ kj/gfw}$, whence he concluded that the primary products of the thermal dissociation are not normal atoms of H and O, but

the temperature.

¹⁹ Hund, F., Z. Physik, 32, 1-19 (1925).

Slater, J. C., Phys. Rev. (2), 38, 1109-1144 (1931).
 Bonhoeffer, K. F., Z. Elektrochem., 34, 652-654 (1928).

²² Gaviola, E., and Wood, R. W., Phil. Mag. (7), 6, 1191-1210 (1928).

²⁸ Senftleben, H., and Reichemeier, O., Physik. Z., 34, 228-230 (1933).

²⁴ Mulliken, R. S., Phys. Rev. (2), 40, 55-62 (1932).

²⁵ Magat, M., Compt. rend., 197, 1216-1220 (1933).

normal H and activated (O1n) oxygen. He stated that this had been previously suggested by Haber and Bonhoeffer.

D. W. Mueller and H. D. Smyth 26 have reported that an electronic bombardment of H₂O-vapor yields negative O-ions and OH-ions, as well as negative H-ions. The number of the H-ions formed per electron is a maximum when the energy of the electrons is about 8 electron-volts; whereas the O-ions and the OH-ions both continue to increase in number with the energy as it exceeds 20 electron-volts.

The energy expended in producing the dissociation accompanying a glow discharge in water-vapor is 11 electron-volts per molecule of H2O dissociated.27

The chemical reactions of water-vapor dissociated by an electrical discharge have been studied by H. C. Urey and G. I. Lavin.²⁸

8 DISSOCIATION OF WATER

Photochemical Dissociation of Water.

When water (liquid) is exposed to the radiation from a quartz-enclosed mercury arc, only H_2 is initially freed, the H_2 O being converted into H_2 O₂; later, from the dissociation of the accumulated H₂O₂, O₂ appears; and finally equilibrium is established, the liberated gas having the composition $2H_2 + O_2$. There is no decomposition of pure water unless the radiation contains waves shorter than 1900A. Between 8 and 20 °C the velocity of decomposition increases as the temperature rises, averaging 1.37 per cent per degree C.16

Ionic Dissociation of Water (See also Electrolytic Ionization, Section 50).

The heat Q_{ton} absorbed in the ionization of water $(H_2O \rightarrow H^+ + OH^-)$ has been expressed by formulas equivalent to the following:

$$(Q_{\text{ion}})_{\text{Ro}} = 57.370 - 0.242(t-18) + 0.00063(t-18)^2 \text{ kj/gfw},$$

 $10 \ge t \ge 35 \,^{\circ}\text{C}^{80};$

$$(Q_{\text{ion}})_{\text{RI}} = 122.16 - 0.2216T \text{ kj/gfw}^{81};$$

$$(Q_{\text{ion}})_{LG} = 57.07 - 0.217(t - 20) \text{ kj/gfw, temperatures near } 20 \,^{\circ}\text{C}^{\,82}$$

$$(Q_{\text{ton}})_{\text{HH}} = 91.760 - 5.9352(T/100) - 1.9862(T/100)^2 \text{ kj/gfw.}^{38}$$

These formulas lead to the values in Table 10.

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26 Mueller, D. W., and Smyth, H. D., Phys. Rev. (2), 38, 1920 (A) (1931).
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²⁷ Linder, E. G., Phys. Rev. (2), 38, 679-692 (1931).

²⁸ Urey, H. C., and Lavin, G. I., J. Am. Chem. Soc., 51, 3290-3293 (1929).

²⁰ Güntherschulze, A., and Schnitger, H., Z. Physik, 103, 627-632 (1936).

²⁰ Rossini, F. D., Bur. Stand. J. Res., 6, 847-856 (RP309) (1931).

Randall, M., Int. Crit. Tables, 7, 232 (1930).
 Lambert, R. H., and Gillespie, L. J., J. Am. Chem. Soc., 53, 2632-9 (1931).

⁸⁸ Harned, H. S., and Hamer, W. J., J. Am. Chem. Soc., 55, 2194-2206 (1933).

Table 10.—Heat of Ionization of Water

		Unit of	Qion = 1	kj/gfw-	H ₂ O; te	$mp. = t \circ ($	$C = T ^{\circ}I$	ζ.		
$_{T}^{t}$	0	10	15	18	20	25	30	40	50	60
4	273.1	283.1	288.1	291.1	293.1	298.1	303.1	313.1	323.1	33 3.1
$(Q_{ion})_{Ro}$ $(Q_{ion})_{Ri}$	61.64	59.35 59.43		57.37 57.65			54.56 54.99	(52.35) 52.78	50.56	48.35
(Qion) LG (Qion) HB	60.74	59.05	(58.16) 58.18	57.50 57.65	57.07 57.30	(55.99) 56.42	51.85	53.71	51.85	49.95

Table 11.—Disposable Energy in the Ionization of Water

The greatest amount of external work that can be obtained from an isothermal ionization of liquid water ($\rm H_2O \rightarrow \rm H^+ + OH^-$) at atmospheric pressure A is $W_{\rm ion} + A\Delta v$; $A\Delta v$ is negligible, $IV_{\rm ion} = -122.16 - 0.2216T \log_e T + 1.40457T$ kj/gfw-H₂O ³⁴; $v_{\rm ion} = W_{\rm ion}/18.0154$. As $W_{\rm ion}$ is negative, work is absorbed during the process. The temperature is t °C, T = 273.1 + t.

	Unit of $W_{ion} = 1 \text{ kj/gfw}$; of $w_{ion} = 1 \text{ kj/g}$.	
t	W_{ton}	wion
0	- 78.07	-4.334
10	-78.72	-4.370
15	- 79.07	- 4.389
20	- 79,44	-4.410
25	-79.82^{a}	-4.431
30	- 80.23	-4.453
40	- 81.10	-4.502
60	- 83.05	-4.610
80	- 85.26	-4.733
100	- 86.90	-4.824

a R. F. Newton and M. G. Bolinger 35 have reported Wion = - 79.92 kj/gfw at 25 °C.

Alpha-ray Dissociation of Water.

The decomposition of water by alpha particles, and the nature of the products formed, have been studied by C. E. Nurnberger.³⁶

⁸⁴ Randall, M., Int. Crit. Tables, 7, 232 (1930).

⁸⁵ Newton, R. F., and Bolinger, M. G., J. Am. Chem. Soc., 52, 921-925 (1930).

⁸⁰ Nurnberger, C. E., J. Phys'l Chem., 38, 47-69 (1934); J. Chem'l Phys., 4, 697-702 (1936).

II. Single-phase Systems

IIA. WATER-VAPOR

9. MOLECULAR DATA FOR WATER-VAPOR

Some miscellaneous data for water-vapor and the numerical values accepted for certain constants appearing throughout this section are given in Table 12, and certain kinetic data for each of a series of temperatures extending from 1 °K to 3000 °C are given in Table 13. More detailed information is given in the following sections and tables. For theory and data consult Boltzmann, 1 Meyer, 2 Jeans, 3 and Dushman. 4

Mean Free Path of Molecules of Water-vapor.

The effective mean free path (L_e) of a molecule of a gas is derived from the observed viscosity (η) by means of the relation $L_e = 3\eta/\rho \bar{v} =$ $3(\pi R/8M)^{1/2} \cdot (\eta T^{1/2}/p) = 3N(\pi/8RM)^{1/2} \cdot (\eta/nT^{1/2})$. When all quantities are expressed in cgs units, this becomes, for water-vapor, $L_e = 4039(\eta T^{\frac{1}{2}}/p) =$ $2.944(\eta/nT^{1/2})\cdot(10^{19})$. Here ρ = density, \bar{v} = mean translational speed of thermal agitation of the molecules, R = gas constant per g-mole, M =molecular weight, p = pressure, $T \circ K = \text{absolute temperature}$, N = number of molecules per g-mole, n = number of molecules per unit of volume.

These relations are derived from the simple kinetic theory of gases devoid of intermolecular attraction. In that case the free path of a molecule between two consecutive collisions is straight, and L_e is a certain average length of all such free paths. In actual gases there is intermolecular attraction, which causes the paths to be curved, increasing the frequency of collision and reducing the lengths of the individual paths. In such cases the value of L_{θ} found by means of the relations just given is the mean free path in a gas having the existing values of M, η , ρ , and \bar{v} , but devoid of intermolecular attraction. It may be called the effective mean free path of the actual gas.

The curvature produced in the path by attraction between the molecules will, obviously, decrease as the translational kinetic energy of the molecules increases. Wherefore W. Sutherland ⁵ suggested that $L_e = L_s/(1 + CT^{-1})$, where L_s is what would be the corresponding mean free path in the actual gas if there were no intermolecular attraction. It may be called the Suther-

² Boltzmann, L., "Vorlesungen über Gastheorie," Leipzig, J. A. Barth, 1896, 1898.

² Meyer, O. E., "Die Kinetische Theorie der Gase," Breslau, Maruschke & Berendt, 1899.

³ Jeans, J. H., "The Dynamical Theory of Gases," Cambridge Univ. Press, 1921.

⁴ Dushman, S., Gen. Elec. Rev., 18, 952-958, 1042-1049, 1159-1168 (1915).

⁵ Sutherland, W., Phil. Mag. (5), 36, 507-531 (1893).

Table 12.-Miscellaneous Molecular Data for Water-vapor

The first three, designated as From the values, R = gas-constant per g-mole = 8.315 10'erg/(g-mole.oK, N = Avogadro's number = number of molecules per g-mole = 6.061×10^{23} , $M = \text{molecular weight of H}_2\text{O} = 18.0154$, and $\eta = \text{viscosity of water-vapor} = 96 \text{ micropoise}$ (= $96 \times 10^{-6} \text{ g/cm·sec}$) at 20 °C, and on the assumption that the molecule of water-vapor is H₂O, the following values are obtained either directly or from the kinetic theory of gases. The first three, designs A, B, and D, respectively, are products that occur frequently in expressions derived from the kinetic theory.

T $^{\circ}$ K.
ø
1 poise = 1 g/cm·sec. Temperature is
g/cm·sec.
poise = 1
; of viscosity = 1
¥
yne/cm2
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Unit of pressure
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Unit

Unit	1 m·sec-1. °K-1/2	10-5 sec. °K 1/2 cm-1	1 (erg/g. °K) *	1 g/g-mole	10^{20}	10' erg/g-mole. 'K	10 ⁻²⁴ g	10-8 g	10-11	10^{18}	1017		1 m/sec	1 m/sec
At 20 °C.	970.2	1.619	2148	18.0154	6061	8.315	29.72	10.85p	4.10p	2.49p	3.65		287	637
At T °K.	370.2	1.619	∞	18.0154	51	15	72	185.7¢T14	$1202.6pT^{-1}$	728.9pT-1	62.4pT-12		34.28714	37.21 T 1/2
Quantity	$A \equiv (2\pi RM)^{1/2}$ (log ₁₀ $A = 4.98684$; cgs units) 970	$= \pi/2A$ (log ₁₀ B = $\overline{5}$.20928; cgs units)	$O \equiv (R/M)^{\frac{1}{2}}$ (log ₁₀ $D = 3.33211$; cgs units) 2148		r = number of molecules per g-mole	Gas-constant per g-mole 8.315	M/N	Mass striking 1 cm ² per sec = $mn_s = p(M/2\pi RT)$ % 185			Number of molecules that strike 1 cm ² of the wall of the container in 1 sec = $n\vec{v}/4 = nP/AT^{12}$ 62.	Mean velocity of the molecules = $D(8T/\pi)^{1/2}$	(See Table 13) 34.2	Square root of the mean squared velocity of the molecules = $D(3T)^{1/4} = \bar{v}(3\pi/8)^{1/4}$ (See also Table 13) 37.2
Symbol	$A = A \equiv$	$B = B \equiv$	$D D \equiv$	M Mol	N Avo.	R Gas-	m Mas	m, Mas	no Nun	" Nun	o Num	v Mea	_	n Squa

E.	Average translational kinetic energy of thermal societion per molecule = $0.5 mm^2 = 1.5 RT/N$			
	(See Table 13)	2.057T	209	10^{-16} erg
E,	As for Em except that E, is referred to the			
	g-mole = $NE_m = 1.5RT$	1.247 <i>T</i>	366	10^{9} ergs
de	Effective diameter of a molecule = $(2KAT^{\frac{1}{2}}/3\pi^{2}N\eta)^{\frac{1}{2}}$			
	(See Molecular size, p. 41).	$104.0(KT\%/\eta)\%$	4.4K%	10-8 cm
d,	Sutherland diameter = $d_0/(1+CT^{-1})^{1/2}$			
	(See Molecular size, p. 41).			
	If $C = 650$ °K, then d_s is	$104.0\{KT^{1.5}/(T+650)\eta\}^{\frac{1}{12}}$ 2.4K%	2.4K%	10^{-8} cm
K	See Table 14 for various values. Preferred value	1.058	1.058	
Γ	Effective mean free path of a molecule			
	$=3\pi/\rho\bar{v}=3NB\pi/nT^{1/2}$ (See also p. 37)	$2.944(10^{\circ}\eta/nT\%)$	16.51/n	$10^{13}\mathrm{cm}$
	Also, $3\eta/\rho\bar{v} = 3D\eta(\pi T/8)^{4h}/p$	4039nT%/p	d/9.9	1 cm
L.	Sutherland mean free path = $L_o(1 + CT^{-1})$			
i	(See p. 37) If $C = 650$ °C	$L_{\bullet}(1+650T^{-1})$	21.7/p	1 cm
nL_s		$29.44(10^{\circ}\eta/T\%)$	165	$10^{12}\mathrm{cm}$
nL.				
İ			531	$10^{12}\mathrm{cm}$
4	Number of collisions per cm ⁸ per sec = $n/\tau = n\bar{v}/L$:		;	•
	$v_{\bullet} = n\bar{v}/L_{\circ} = 8Nb^2/3\pi R\eta T$	$6.19p^2/\eta T$	$219p^2$	10^{15}
•	Moss free time = $I/\bar{m} \cdot \tau_s = I_s/\bar{m} = 3\pi n/8b$	11784/p	0.113/p	10- sec
- 1	Viscosity of water-vapor Value at 20 °C is taken as		96	10-6 g/cm·sec
'n	VISCOSILY OF WALLS TREES.			

land mean free path. The "Sutherland constant" (C) is characteristic of the gas, and measures the intensity of the intermolecular attraction. This suggestion of Sutherland's leads to the relation $\eta = AT^{1.5}/(T+C)$, where $A \equiv nL_8(8RM)^{12}/3N\pi^{1/2}$ is a constant. This relation agrees quite closely with the observations on many gases over wide ranges in T, the pressure not exceeding a few atmospheres. But as the pressure increases, the isothermal value of η for water-vapor increases (see Section 11), and the Sutherland relation ceases to hold. This means that nL_8 is not a constant, as the simple theory demands, but varies with the pressure, the variation depending upon T.

Unless qualified in some way, the term "mean free path" is ambiguous. In common practice, it usually means either L_{θ} or L_{θ} . For water-vapor

Table 13.-Kinetic Data for Molecules of Water-vapor

Information regarding the symbols not defined here and the numerical values of the basic constants will be found in Table 12.

If $\rho =$ density, $p = \frac{1}{3}\rho(v_2)^2 = RTn_0$; whence $(v_2)^2 = 3RT/M$. If the distribution of velocities is Maxwellian, $\bar{v} = v_2\sqrt{8/3\pi} = 0.9213 v_2$. Insofar as the principle of equipartition of energy applies, each g-mole of a resting gas, containing only one type of molecule and in thermal equilibrium, contains an amount of kinetic energy equal to 0.5RT for each of the degrees of freedom characteristic of the molecule.

m= mass of one molecule of $H_2O=29.72\ 10^{-24} g$; $(v_2)^2=1.3846T\ 10^7 (\text{cm/sec})^2$; $v_2=3721.1 \sqrt{T}\ \text{cm/sec}$; $\bar{v}=3428.2 \sqrt{T}\ \text{cm/sec}$; $E_m=$ average translational kinetic energy of thermal agitation of a molecule = $20.57T\ 10^{-17} \text{erg}$; $E_g=$ the same kinetic energy of a g-mole = $NE_m=12.472T\ 10^7$ ergs.

Unit of \vec{v}_i and $\vec{v}_i = 1$ m/sec of $E_m = 10^{-16}$ ergs of E_n and of $0.5RT = 10^8$ ergs.

Om	tot v and vg	I mysec, or 2	10 016,	or Ly and or	0.5111 - 10	C. 85.
ŧ	T	$ar{v}$	v_2	E_{m}	E_{g}	0.5RT
-272.1	1	34.3	37.2	2.1	1.2	0.42
-271.1	2	48.5	52.6	4.1	2.5	0.83
-268.1	5	76.6	83.2	10.3	6.2	2.08
-263.1	10	108.4	117.7	20.6	12.5	4.16
-250	23.1	164.8	178.8	47.5	28.8	9.60
-200	73.1	293.1	318.1	150.4	91.2	30.39
-150	123.1	349.5	379.4	253.2	153.5	51.18
-100	173.1	451.0	489.6	356.1	215.9	71.97
-50	223.1	512.0	555.8	458.9	278.2	92.7
O	273.1	566.5	614.9	562.	340 .6	113.5
25	298.1	591.9	642.5	613.	371.8	123.9
100	373.1	662.2	718.8	767.	465.3	155.1
200	473.1	745.7	809.4	973.	590.0	196.7
300	573.1	820.7	890.8	11 7 9.	715.	238.3
500	773.1	953.2	1034.6	1590.	964.	321.4
1000	1273.1	1223.2	1327.7	2619.	1588.	529.3
1500	1773.1	1443.5	1566.9	3647.	2211.	737.2
2000	2273.1	1634.5	1774.1	4676.	<i>2</i> 835.	945.0
3000	3273.1	1961.3	2128.9	6733.	4082.	1360.8

 $nL_{\rm e}$ is about 1.6×10^{14} cm at room temperature and saturation pressure, and increases with both pressure and temperature to about twice that value at the limits of the experimental values for the viscosity; the corresponding values of $nL_{\rm e}$ are about 5 and 8×10^{14} cm, varying not greatly with T but subject to variations in the chosen value of C (see Table 21). At a pressure of 1 barye (1 dyne/cm²) $L_{\rm e}$ varies from about 6 cm at 0 °C to about 20 cm at 350 °C, the corresponding values of $L_{\rm e}$ being 20 and 45 cm respectively.

Molecular Size.

A general discussion of the various methods available for estimating the sizes of molecules, atoms and ions, together with numerical values and a bibliography of 153 titles has been published by K. F. Herzfeld.⁶

In the simple kinetic theory of gases, the term *molecular diameter* is used to indicate the diameter of the equivalent elastic spheres by which the molecules may be regarded as replaced when one is concerned solely with the mechanical effects arising from the thermal agitation of the molecules of the gas when in a state closely approximating the ideal. This diameter is an average minimum value of the distance between the centers of colliding molecules under the stated conditions. It is an average, because the minimum distance may vary from one collision to another, because the molecules have various velocities, are neither necessarily spherical nor surrounded by fields having spherical symmetry, and possibly interpenetrate, more or less, at each collision.

Insofar as the molecules may be replaced by elastic spheres between which there are no forces except when the spheres are in actual collision, the path between two consecutive collisions will be straight, and the average value (L) of all such paths will be related to d, the diameter of the spheres, as indicated by the equation $nL = K/\pi d^2$, n being the number of molecules per unit volume and K being a number determined by the distribution of the velocities of the molecules and by the kind of average free path represented by L. Some of the values that have been computed for K are given in Table 14.

It seems to be agreed that Chapman's highest value (K=1.058) is the most appropriate for use in formulas involving the viscosity. But all the other values tabulated have been used, and molecular diameters so computed are not infrequently tabulated without any indication of the value of K to which they refer, even though different entries refer to different values, and some refer to $d_{\mathfrak{e}}$ (computed from $L_{\mathfrak{e}}$) and some to $d_{\mathfrak{e}}$ (computed from $L_{\mathfrak{e}}$).

The area πd^2 is often called the cross-sectional area of the sphere of action. But when that term is used, d having the significance here considered, it should be so qualified as to indicate that it refers solely to those intermolecular collisions that occur as the result of thermal agitation; for

e Herzfeld, K. F., Jahrb. d. Radioak., 19, 259-334 (1922).

the radius of the sphere of action may have a different value in relation to other events. From the observed viscosity of water-vapor it may be inferred that d_c is of the order of 3 to 5A (1A = 10^{-8} cm), and d_s of 2A; the corresponding values of πd^2 are of the orders of 28, 78, and $13A^2$, respectively.

Table 14.—Various Values Assumed for $K =$	$\pi d^2 nL$
--	--------------

K is a pure number.	K
I. Gas at rest; $L = \text{mean value of the free paths}$:	
(a) All molecules, except one, continuously at rest	1.
(b) All molecules have the same speed, chaotically distributed	0.750
(c) Maxwellian distribution of velocities $K = 1/\sqrt{2}$	0.707
 II. Gas streaming lamellarly; various types of averaging, each intended to be that appropriate to the equation η = ½ρēL: (a) Distribution of velocities strictly Maxwellian. K = 	
$3(0.3502)/\sqrt{2} = 1.0506/\sqrt{2}$	0.743
(b) Maxwellian distribution as modified by the streaming ² $K = 3(0.3097)/\sqrt{2} = 0.9290/\sqrt{2}$	0.657
(c) Taking into account the persistence of the streaming velocity (it was ignored by Boltzmann and by Meyer), and assuming a Maxwellian distribution as modified by streaming, J. H. Jeans 3 finds by one approximation $K=1.317/\sqrt{2}\ldots$ and by another $K=1.382/\sqrt{2}\ldots$	0.931 0.977
(d) By a somewhat different procedure, S. Chapman ⁷ finds for condition (c) $K = 3(0.491)(1 + \epsilon)/\sqrt{2} = 1.473$ $(1 + \epsilon)/\sqrt{2} = 1.042(1 + \epsilon)$, where $(1 + \epsilon)$ depends upon the intermolecular attraction, and lies between 1.000 and 1.016. If $\epsilon = 0$, K is	1.042 1.058

Unfortunately, the term *molecular diameter* is not infrequently used in other senses. This has led to great confusion, and not infrequently to comparison of values that are not comparable. Some of these other uses are illustrated in Table 15. Each of the values there tabulated has been called molecular radius. Some have been called radius of molecular action, and some half the radius of molecular action.

⁷ Chapman, C., Phil. Trans., (A), 216, 279-348 (1916).

Table 15.—Estimates of the Effective Size of the Molecule of Water-vapor

r= "radius" of the molecule as derived in the manner indicated. If the procedure defines a diameter (d), then r=d/2. For additional explanations, see text.

•	
Unit of $r = 1A - 10^{-8}$ cm. Basis	,
Viscosity at 100 °C and 1 atm; $r = d_c/2$	2.08
Viscosity at 100 °C and 1 atm; $r = d_s/2$	1.25
Van der Waals' equation, assuming that the co-volume b is 4 times the volume of all the molecules in a g-mole, that the critical pressure and temperature are $P_c = 217.7$ atm. and $T_c = 273 + 374$ °C = 647 °K, and that $b = RT_c/8P_o$	1.44
As before, except that $b = RT_c/15P_c$	1.17
Kamerlingh-Onnes' equation of state and Jakob's data for the density. G. Holst ⁸ finds	3.2
The relation $\pi d^3 n/6 = (\epsilon - 1)/(\epsilon + 2)$, $\epsilon = \text{dielectric constant}$	2.9
If in the liquid state the molecules were cubes and were closely packed at 4 °C, then the length of the edge of each cube would be $c = (M/N\rho)^{1/3} = 2r$	1.55
If in the liquid state the molecules were spheres and were closely packed at $4 ^{\circ}\text{C}$ (tetrahedral packing), then $d = c \cdot 2^{1/6} = 2r \dots$	1.74
If δ is mean distance between the centers of adjacent molecules of a liquid in which the velocity of sound is V and the thermal conductivity is λ , then $\delta = (2kV/\lambda)^{\frac{1}{2}} = 2r$, where k is the Boltzmann gas constant $= R/N = 1.372 \times 10^{-16} \text{ (erg/°K)}$ per molecule. For water at $4 ^{\circ}\text{C}$, $\lambda = 5.61 \times 10^4 \text{erg/cm·sec·°C}$, $V = 1425 \text{m/sec}$.	1.32
Assuming that the effective radius of a molecule of mercury vapor is 1.80A (the value computed from the viscosity, but whether in terms of L_c or of L_s , and if the latter, for what temperature, is not stated, M. W. Zemansky 9 determined the greatest value of $r = \Delta - 1.80$ A that is consistent with the production of a stated effect upon the radiation emitted by the mercury molecule, Δ being the distance between the center of a water-vapor molecule and that of a colliding molecule of mercury vapor. His data yield the following values:	
For quenching the radiation	
For depolarizing the radiation	
For broadening the spectral lines	6.48
8 TI -1-4 C D Ab -1 Tiles Americanism 10 022 027 (1017)	

⁸ Holst, G., Proc. Akad. Wct. Amsterdam, 19, 932-937 (1917).

^o Zemansky, M. W., Phys. Rev. (2), 36, 919-934 (1930).

Table 15—(Continued)

Basis

Dasis	
If Δ is the greatest mean distance between the center of a molecule	
of water-vapor and that of an electron, consistent with the mole-	
cule's essential blocking of the advance of the electron, then the	
value of Δ for each of several values of the kinetic energy of the	
colliding electron can be determined from a curve published by	
E. Brüche. ¹⁰ They are as follows, Δ being taken as r :	
Energy = 4 electron volts	2.24
Energy = 7.8 electron·volts	2.32
Energy = 9 electron volts	2.39
Energy = 16 electron volts	2.30
Energy = 25 electron volts	2.03
Energy = 36 electron volts	1.83
Formulas connecting the diameter of a molecule of a vapor with the	
surface tension and other properties of the corresponding liquid	
have been derived from assumptions that have not been generally	
accepted. Thus:	
S. Mokroushin 11 obtains a value, which after correction to	
the basis of 6.06×10^{23} molecules per g-mole, yields	2.72
S. P. Owen 12 obtains	1.62
In the triangular molecule (Table 16) the distance from the center	
of the oxygen to that of either hydrogen nucleus is approximately	1.00

Moments of Inertia of the Molecule of Water-vapor.

Band spectra are attributed to the rotation of the molecule. The fundamental frequencies, from which those of the maxima of the several bands are obtained by summations and differences, are determined by the frequency of rotation of the molecule, as are also the constant frequencydifferences between the consecutive lines of a band and between the components of certain doublets. The quantum theory establishes a relation between these frequencies, or differences in frequency, and the changes in the rotational energy of the molecule. From that, the pertinent moment of inertia of the molecule can be obtained.

For example, ignoring all complications, the quantum theory requires that the integral of the angular momentum over a complete cycle shall be an integral (n) multiple of the Planck constant of action ($h = 6.56 \, 10^{-27}$ erg·sec); that is, $4\pi^2 I \bar{\nu} = nh$, I being the effective moment of inertia, and $\bar{\nu}$ the frequency of the rotation of the molecule. Hence, the rotational

energy is $E = \frac{1}{2}I(2\pi\bar{\nu})^2 = \frac{1}{8\pi^2I}(h^2n^2)$. As *n* changes from one integer

Brüche, E., Ann. d. Physik, (5), 1, 93-134 (1929).
 Mokroushin, S., Phil. Mag. (6), 48, 765-768 (1914).

¹² Owen, S. P., Proc. Univ. Durham Phil. Soc., 6, 308-311 (1932).

to the next, E changes by $\Delta E = \frac{h^2}{8\pi^2 I} (2n+1)$; and on the quantum

theory, this must equal $h\nu$, where ν is the frequency of the associated radiation. Whence it is evident that each unit change in n causes ν to change by $\Delta \nu = h/4\pi^2 I$, which gives rise to a spectrum of lines spaced at equal intervals of frequency. From the observed values of $\Delta \nu$, I can be computed:

$$I = h/4\pi^2 \Delta \nu = 1.66_1/\Delta \nu = 55.4/(1/\lambda_1 - 1/\lambda_2) 10^{-40} \text{ g} \cdot \text{cm}^2$$

the unit of λ being 1 cm, and of ν being 1 wave/sec. In other cases the procedure is somewhat similar. In every case I is the effective moment of inertia under the conditions characteristic of the vibrations by means of which it has been obtained; it differs from that pertaining to the static molecule.^{13, 14} An early review of the subject was published by A. Eucken,¹⁵ and a later one may be found in a monograph by C. Schaefer and F. Matossi.¹⁶

The absorption spectrum of water-vapor is so complex that until recently its interpretation has been very incomplete and subject to dispute, but the difficulties are now being rapidly overcome. Three different moments of inertia are involved, indicating that the atoms in a molecule lie at the vertices of a triangle.^{13, 14, 15; 16, p. 235-245; 17, 18, 19, 20, 21} The values of the moments of inertia as derived by F. Hund ²² from the observations by Eucken ¹⁵ have been much quoted and used; they are $I_A = 0.98$, $I_B = 2.25$, $I_C = 3.20 \ 10^{-40} \text{g·cm}^2$, or $I_A = 0.59$, $I_B = 1.34$, $I_C = 1.91$ protonangstrom.² But probably the best values are those given by K. Freudenberg and R. Mecke ¹⁴ and based on the structure of 17 water-vapor bands studied by Mecke and his associates. They are as follows, the unit being $10^{40} \text{ g·cm}^2 = 0.5969 \ m_{\text{H}} A^2$, m_{H} indicating the mass of a hydrogen atom:

$$I_1 = 0.996 + 0.045\sigma + 0.026\pi - 0.0988$$

 $I_2 = 1.908 + 0.014\sigma + 0.033\pi - 0.0348$
 $I_3 = 2.981 + 0.047\sigma + 0.062\pi + 0.0628$
 $\Delta = 0.077 - 0.012\sigma + 0.003\pi + 0.1948$

where $\Delta \equiv I_8 - (I_1 + I_2)$, and (σ, π, δ) are the quantum integers involved in the vibration considered. For the stationary (unvibrating) state they give $I_1 = 1.009$, $I_2 = 1.901$, $I_3 = 2.908$.

The values found by P. Lueg and K. Hedfeld ²¹ from a study of 3 bands are $I_A = 0.97$, $I_B = 2.13$, $I_C = 3.07$, but R. Mecke and W. Baumann ¹³ think that the data they used are not satisfactory.

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    Mecke, R., and Baumann, W., Physik. Z., 33, 833-835 (1932).
    Freudenberg, K., and Mecke, R., Z. Physik, 81, 465-481 (1933).
    Eucken, A., Jahrb. d. Radioak., 16, 361-411 (1920).
    Schaefer, C., and Matossi, F., "Das Ultrarote Spektrum," Berlin, Julius Springer, 1930.
    Witt, H., Z. Physik, 28, 249-255 (1924).
    Mecke, R., Physik. Z., 30, 907-910 (1929).
    Mahanti, P. C., Physik, Z., 32, 108-110 (1931).
    Mecke, R., Z. Physik, 81, 313-331 (1933).
    Lueg, P., and Hedfeld, K., Z. Physik, 75, 512-520 (1932).
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From the data available in 1927, R. T. Birge 23 derived the following values for the OH-ion, h being taken as 6.557 10 27 erg-sec and c as 2.99796 10^{10} cm/sec; moment of inertia = 1.634 10^{40} g·cm² at the upper state of excitation, and 1.500 at the lower, the corresponding separation of the nuclei being 1.022A and 0.979A. On the same basis, but from more recent data, D. Jack 24 derived essentially the same values for the moments of inertia, *i.e.*, 1.633 and 1.498 10^{-40} g·cm². While from their own measurements of a newly discovered band, H. L. Johnston, D. H. Dawson, and M. K. Walker 25 derive for the states 2_{Σ} and 2_{π} , respectively, the values $10^{40}I = 1.591$ and 1.454 g·cm², and the separations 1.009A and 0.964A.

A review of the methods available for estimating the moments of inertia and certain related data has been published by A. Eucken ¹⁵; more detailed treatments may be found in various treatises, such as C. Schaefer and F. Matossi's "Das Ultrarote Spectrum." The derivation of formulas required for the interpretation of the spectrum has been published by F. Lütgemeier, ²⁶ II. A. Kramers and G. P. Ittmann, ²⁷ D. M. Dennison, ²⁸ and others.

Dipole Moment of the Molecule of Water-vapor.

Several distinct phenomena exhibited by some substances, but not by others, can be most satisfactorily explained by assuming that the molecules involved in them contain rigid, or nearly rigid, electrical dipoles, the distance between the poles being significantly less than the diameter of the molecule. Water-vapor exhibits phenomena that can be explained in this way. Discussions of the general subject have been published by P. Debye ²⁹; and by the Faraday Society,³⁰ and of certain aspects of it by R. S. Mulliken.³¹

In 1903, M. Reinganum ³² attributed intermolecular forces to such dipoles, and on that assumption computed the moments that must be assigned to them for each of a number of liquids. But the magnitude of these moments can be obtained more directly from a study of the dielectric constants, as has been pointed out by P. Debye.³³

He showed that if each molecule contains a rigid electrical dipole as

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<sup>22</sup> Hund, F., Z. Physik, 31, 81-106 (1925).

<sup>23</sup> Birge, R. T., Int. Crit. Tables, 5, 415 (1929).

<sup>24</sup> Jack, D., Proc. Roy. Soc. (London) (A), 118, 647-654 (1928).

<sup>25</sup> Johnston, H. L., Dawson, D. H., and Walker, M. K., Phys. Rev. (2), 43, 473-480 (1933).

<sup>26</sup> Lütgemeier, F., Z. Physik, 38, 251-263 (1926).

<sup>27</sup> Kramers, H. A., and Ittmann, G. P., Idem, 53, 553-565 (1929); 58, 217-231 (1929); 60, 663-681 (1930).

<sup>28</sup> Dennison, D. M., Rev. Mod. Physics, 3, 280-345 (1931).

<sup>29</sup> Debye, P., "Handb. d. Radiol." (E. Mark. Ed.), 6, 597-786 (1925); "Polar Molecules," New York Chemical Catalogue Co. (Reinhold Publishing Corp.), 1929.

<sup>30</sup> The Faraday Society, Trans., 30, 679-904 (1934).

<sup>31</sup> Mulliken, R. S., J. Chem'l Phys., 3, 573-585 (1935).

<sup>32</sup> Reinganum, M., Ann. d. Physik (4), 10, 334-353 (1903).

<sup>33</sup> Debye, P., Physik. Z., 13, 97-100, 295 (1912).

<sup>34</sup> Debye, P., "Polar Molecules," p. 8, New York, Chemical Catalog Co. (Reinhold Publishing Corp.), 1929.
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well as electrons that are elastically bound, relation (1) should be satisfied,

$$\frac{\epsilon - 1}{\epsilon + 2} \cdot \frac{1}{\rho} = a + \frac{b}{T} \tag{1}$$

the intermolecular action of the fields due to the molecules themselves being considered to a first approximation only. In this equation, $\epsilon = \text{dielectric}$ constant, $\rho =$ density, a and b are essentially positive constants characteristic of the substance, and T is the absolute temperature. The first term (a)depends upon the elastically bound electrons and determines the optical dispersion of the substance; the second constant (b) depends upon the strength, or moment, of the dipole. If the electrical quantities are expressed

in electrostatic units, then $a = \frac{4\pi}{3} \cdot \frac{N}{M} \alpha$ and $b = \frac{4\pi \hat{N}}{9kM} \mu^2$, where N = num

ber of molecules per mole, M = molecular weight, k = molecular gasconstant = R/N, R being the gas-constant per mole, μ = moment of the dipole, and α the moment induced by unit field. The quantity α is called the "polarizability" 34 or the "deformability" 35 of the molecule. $\alpha \equiv$ $e^2\Sigma_p(n/f)_p$ where e= electronic charge, $n_p=$ number, per molecule, of the elastically bound electrons for which the force of restitution is f_p per unit displacement, and Σ_p = summation for all values of p. When in an electrostatic field of strength E, such a molecule will have an induced electric moment of amount $\mu' = \alpha E$. The value of a is related to the index of refraction (n) and to the density (ρ) as follows: $a\rho = (n^2 - 1)/(n^2 + 2)$.

Taking the unit of $\rho = 1$ g/cm³, $N = 6.06_1 \times 10^{23}$ molecules per g-mole, M = 18.0154, $k = 1.372 \, 10^{-16}$ erg per molecule and per °K(I.C.T. values), we find for water-vapor:

$$10^{24} \alpha = 7.10a \text{ cgsc units}$$

 $10^{20} \mu = 5.40 \sqrt{b} \text{ cgsc units}$

R. Gans ³⁶ has considered in greater detail the effect of the intermolecular fields upon ϵ ; and convenient methods for using the more complicated equations so obtained have been given by H. Isnardi,37 P. Lertes,38 and C. P. Smyth.³⁹ But the simpler formulas given by Debye are amply accurate for our present purposes.

The available data for water-vapor are shown in Figure 2, Section 22, $(\epsilon - 1)T/\rho$ being plotted against T. As $(\epsilon + 2)$ is practically equal to 3, equation (1) requires that the points shall lie on a right line, sloping upward as T increases, and cutting the axis of ordinates at $(\epsilon - 1)T/\rho = 3b$. The data published by J. D. Stranathan 40 seem to be by far the most satis-

⁸⁵ Rao, I. R., Indian J. Phys. 2, 435-465 (1928).

³⁸ Gans, R.. Ann. d. Physik (4), 64, 481-512 (1921).

³⁷ Isnardi, H., Physik. Z., 22, 230-233 (1921).

⁸⁸ Leites, P., Z. Physik, 6, 257-268 (1921).

⁸⁹ Smyth, C. P., Phil. Mag. (6), 45, 849-864 (1923); J. Am. Chem. Soc., 46, 2151-2166 (1924). 40 Stranathan, J. D., Phys. Rev. (2), 48, 538-549 (1935) → 47, 794(A) (1935), extending 45, 741 (1934).

factory. They lead to $b = 1149.6 \pm 7.8$, giving

 $10^{18} \mu = 1.83_1 \text{ cgse}$ $\mu = 0.383$ electron angstrom

Another recent set of data is that by L. G. Groves and S. Sugden, 41 leading to $10^{18} \mu = 1.84 \pm 0.01$ cgse.

Of earlier data, the most satisfactory set is that published by R. Sänger and O. Steiger 42; but the published details are not sufficient to enable one to form an independent estimate of the accuracy of the final results. They lead to a value of 10¹⁸ µ lying between 1.84 and 1.88 cgse, a little greater than Stranathan's value.

The observations of C. T. Zahn 48 are not inconsistent with these values for μ , but are too scattered to justify an attempt to derive an independent value from them. The same is true of M. Jona's,44 which are admittedly affected by an unknown percentile error.

The data of K. Bädcker, 45 though frequently quoted, and used by G. Holst 46 in his computation of u, are plainly in error and are worthless for that purpose, as they impose a negative value upon the essentially positive a.

Less directly obtained estimates have been made, based upon more or less questionable assumptions and in some cases involving rough empirical relations. Thus, G. Holst 46 has computed from Kamerlingh-Onnes' equation of state, as fitted to Jakob's data for the density, the value $10^{18} \mu = 2.6$ cgse; whereas J. K. Syrkin ⁴⁷ has concluded that $10^{20} \mu = 1.66 T_{\rm crit} / (P_{\rm crit})^{1/2}$, the unit of P being 1 atm, thus finding for water-vapor $10^{18} \mu = 0.73$ cgse. (Syrkin does not state what units he uses, but they seem to be as here given; however, certain of the relations that he gives appear to be mutually inconsistent, no matter what probable guess is made regarding the units.)

A. Kirrmann 48 has reviewed the various methods that have been proposed for estimating the value of μ , giving numerical values for various substances and a bibliography of 42 entries. Numerous errors occur in both tabulation and bibliography, and in at least some instances data derived from observations on the liquid phase by procedures now known to be incorrect are not distinguished from those derived satisfactorily from observations on the corresponding gas phase. A more recent table of dipole

⁴¹ Groves, L. G., and Sugden, S., J. Chem. Soc. (London), 1935, 971-974 (1935). ⁴² Sänger, R., and Steiger, O., *Helv. Phys. Acta.* 1, 369-384 (1928); republished by Sänger, R., *Physik. Z.* 31, 306-315 (1930), and by Sänger, R., Steiger, O., and Gächter, K., *Helv. Phys. Acta.* 5, 200-210 (1932).

⁴⁸ Zahn, C. T., Phys. Rev. (2), 27, 329-340 (1926).

⁴⁴ Jona, M., Physik. Z., 20, 14-21 (1919). 45 Bädeker, K., Z. physik. Chem., 36, 305-335 (1901).

⁴⁶ Holst, G., Proc. Akad. Wet. Amsterdam, 19, 932-937 (1917).

⁴⁷ Syrkin, J. K., Z. anorg. allgem. Chem., 174, 47-56 (1928).

⁴⁸ Kirrmann, A., Rev. gén. des Sci., 39, 598-603 (1928).

moments, collected by N. V. Sidgwick, has been published by G. C. Hampson and R. J. G. Marsden.49

Polarizability of the Molecule of Water-vapor.

The term "polarizability" of the molecule is used by Debye to denote that portion of the molecular electric moment induced per unit electrical field as a result of the displacement of elastically bound electrons. It is the quantity already denoted by α , and is related to the α of equation (1) as follows: $a = (4\pi N/3M)\alpha$, which in the case of water-vapor becomes $10^{24} \alpha = 7.10a$ cgse units (see preceding topic: Dipole Moment). Both Raman and Rao call it the (mean) "deformability" of the molecule. The quantity a is related to the index of refraction (n) and the density (ρ) in this manner: $a_{\rho} = (n^2 - 1)/(n^2 + 2)$.

From the last stated relation, I. R. Rao 85 has computed the value $10^{24} \alpha = 1.50$ cgse. From the α (0.224) derived from Stranathan's observations (1935) (Section 22), we find

> $10^{24} \alpha = 1.59 \text{ cgse}$ $\alpha = 1.59$ electron angstrom per (e/A^2) field $= 1.59A^3$

That is, if a molecule of water-vapor is placed in an electro-static field having n times the strength of the field that exists at the distance of 1 angstrom (10⁻⁸ cm) from an isolated electron (charge = $e = 4.774 \times 10^{-10}$ cgse), then that field will induce in the molecule an electric moment equivalent to that of two charges of opposite signs, each equal to e, separated by the distance of 1.59n angstroms, n being assumed to be so small that the induced moment is sensibly proportional to the strength of the inducing field.

This polarizability is generally ascribed almost exclusively to the oxygen. The polarization of the oxygen by its attendant hydrogens gives rise to a very significant negative component in the permanent dipole moment of the water-vapor molecule. See P. Debye, 34, p. 68+ and I. R. Rao. 35

Anisotropy of the Molecule of Water-vapor.

Light scattered at an angle of 90° by water-vapor, and by certain other gases and vapors, is not completely polarized. This suggests that the polarization induced in the molecule by the incident light depends upon the orientation of the molecule, the latter being optically anisotropic. Similarly, any existence of electric or of magnetic double refraction would indicate that the molecule is anisotropic with reference to those forces also. For theoretical treatment of the subject, see T. H. Havelock 50; C. V. Raman and K. S. Krishnan, 51 F. Hund, 52 and P. Debye. 53

Hampson, G. C., and Marsden, R. J. G., Trans. Faraday Soc., 30, appendix, 86 pp. (1934).
 Havelock, T. H., Phil. Mag. (7), 3, 158-176 (1927).
 Raman, C. V., and Krishnan, K. S., Idem, 713-723, 724-735 (1927).

If the electrical moments induced along the principal axes of the molecule by unit external fields parallel to those axes are A, B, and C, then, in the notation of Raman and of Rao

 δ = factor measuring the anisotropy of the molecule =

$$\frac{A^2 + B^2 + C^2 - (AB + BC + CA)}{(A + B + C)^2}$$

r = depolarization factor = ratio of the weaker plane-polarized component of the transversely scattered light to the stronger one.

A, B, C = deformabilities of the molecule.

a, b, c = deformabilities of the atom or ion.

 $\alpha=$ (mean) deformability of the molecule $=\frac{3}{4\pi N_1}\left(\frac{n^2-1}{n^2+1}\right)$, n= index of refraction, $N_1=$ number of molecules per unit volume. Debye calls α the polarizability.

When magnetic anisotropy is to be considered, the components of the susceptibility (or magnetic deformability) are referred to the same axes as A, B, and C; and are commonly denoted by A', B', and C'. Similarly in other cases.

For water-vapor, B = C, approximately; whence, writing $\Delta \equiv A/C$, $\delta =$

$$\left(\frac{\Delta-1}{\Delta+2}\right)^2; \ r=\frac{2(\Delta-1)^2}{4(\Delta-1)^2+5(2\Delta+1)}; \ \alpha=\frac{A+2C}{3}=C\left(\frac{\Delta+2}{3}\right).$$

I. R. Rao ⁵⁴ found r=0.0191, $r_0=0.0199$; $\delta=0.0166$ at 120 °C, r_0 being the value corresponding to a very low density. In a later paper ⁵⁵ Rao suggested that the atoms and ions may themselves be anisotropic, and concluded that the anisotropy of O⁻⁻ is greater in H₂O than in CO₂, and "seems to be anomalous." He attributed the entire anisotropy of H₂O to the O⁻, and gave for it $\Delta=A/C=a/c=1.45$ to 1.53.

All of this refers to the optical anisotropy of the molecule. There appears to be no available data from which the magnetic anisotropy of a molecule of water-vapor can be computed.

It should be remarked that the notation in this field is confused. For example, S. W. Chinchalkar ⁵⁶ defines the optical anisotropy as

$$\delta = \frac{(A-B)^2 + (B-C)^2 + (C-A)^2}{(A+B+C)^2}$$

which is twice the value used by Raman and Rao.

⁵² Hund, F., Z. Physik, 43, 805-826 (1927).

⁵⁸ Debye, P., "Handb. d. Radiologie" (E. Marx, Ed.), 6, 754-776, Leipzig, 1925.

⁵⁴ Rao, I. R., Indian J. Phys., 2, 61-96 (1928).

⁵⁸ Rao, I. R., Idem, 2, 435-465 (1928).

⁵⁶ Chinchalkar, S. W., Indian J. Phys., 6, 165-179 (1931).

Models of the Molecule of Water-vapor.

Descriptive models of the molecule have been derived from chemical data. These need not detain us beyond mentioning that C. Friedel,⁵⁷ J. W. Brühl,⁵⁸ H. E. Armstrong,⁵⁹ T. M. Lowry and H. Burgess,⁶⁰ J. Piccard,⁶¹ C. P. Smyth,⁶² M. Smith,⁶³ M. L. Huggins,⁶⁴ and others have presented reasons for supposing that oxygen is tetravalent, that the electrons of the oxygen atom have a tetrahedral distribution about the nucleus, and that the H₂O-molecule is a tetrahedron, two of its vertices being occupied by H and two by electrons serving as bonds for secondary valencies.

On this basis, S. W. Pennycuick 65 has constructed models for ice and for water.

Another mode of approach is by way of physical data. This leads to numerical results and to an idea of the probable stability of the model proposed. Only two types of model are possible. Either the three molecules lie in a straight line, or they do not; the first gives a linear, and the second a triangular model. The latter is in qualitative accord with that mentioned in a preceding paragraph.

It may be shown that the linear model is either non-polar or unstable.^{34, pp. 63-68} Hence this model, which was considered by T. H. Havelock ⁵⁰ and proposed by F. J. v. Wisniewski, ⁶⁶ must be condemned, as we know that the H₂O-molecule is polar (see *Dipole Moment*, p. 46).

The triangular model is stable ^{34, pp. 68-76} if the triangle is isosceles, the oxygen being at the unique vertex, and if the oxygen is so polarizable that $\alpha/r^3 > 1/8$, where r is the distance from the nucleus of the oxygen atom to that of either hydrogen atom, and α is the polarizability (p. 47). Furthermore, if the unique internal angle of the triangle is 2θ , $\alpha = (1/8)(r/\sin\theta)^3 = (r^2/b)^3$ and the total electric moment of the molecule is $\mu = 2er\cos\theta(1 - 1/8\sin^3\theta) = 2ca(1 - r^3/b^3)$, where a is the altitude of the triangle, and b the base.

Such a triangular arrangement has been discussed by W. Heisenberg, 67 M. Born and W. Heisenberg, 68 F. Hund, 22 T. H. Havelock, 50 P. Debye, $^{34, pp. 63-68}$ A. S. Coolidge, 69 and others. Regarded as a rigid body, it has three principal moments of inertia: I_a about the axis along a, I_b about that parallel to b and passing through the center of mass of the system, and I_a about the axis perpendicular to the plane of the three atoms

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57 Friedel, C., Bull. Soc. Chim. de France (N.S.), 24, 160-169, 241-250 (1875).

18 Bruhl, J. W., Ber. deut. chem. Ges., 28, 2866-2868 (1895) ← Z. physik. Chem., 18, 514-518 (1895); Ber. deut. chem. Ges., 30, 162-172 (1897).

18 Armstrong, H. E., Compt. rend., 176, 1892-1894 (1923).

19 Lowry, T. M., and Burgess, H., J. Chem. Soc. (London), 123, 2111-2124 (1923).

20 Enyth, C. P., Phil. Mag. (6), 47, 530-544 (1924).

21 Smyth, C. P., Phil. Mag. (6), 47, 530-544 (1924).

22 Smyth, M., "Chemistry and Atomic Structure," 1924.

23 Huggins, M. L., Phys. Rev. (2), 27, 286-297 (1926).

24 Pennycuick, S. W., J. Phys'l Chem., 32, 1681-1696 (1928).

25 W. Wisniewski, F. J., Z. Physik, 47, 557-588 (1928).

26 Heisenberg, W., Z. Physik, 26, 196-204 (1924).
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** Born, M., and Heisenberg, W., Idem, 23, 388-410 (1924). ** Coolidge, A. S., Phys. Rev. (2), 42, 187-209 (1932).

and passing through the center of mass. If M = mass of an oxygen atom = 26.40 10^{-24} g, m = that of a hydrogen atom = $1.676 \ 10^{-24}$ g, and a = mass and b = mass are expressed in centimeters, then $I_a \equiv 0.5 mb^2 = 0.838b^2 \ 10^{-24} \text{g·cm}^2$; $I_b \equiv 2Mm \ a^2/(M+2m) = 2.974a^2 \ 10^{-24} \text{g·cm}^2$; and $I_a = I_a + I_b$. Whence, $10^{-12}a = 0.580(I_b)^{0.5}$, $10^{-12}b = 1.092(I_a)^{0.5}$, $\tan \theta = b/2a = 0.942 \ (I_a/I_b)^{0.5}$.

Spectral data show that the molecule H_2O has indeed three principal moments of inertia, and furnish the means for determining their values (Tables 64 and 65). The largest is evidently I_o , but it is not obvious which of the others should be assigned to I_a . Of the two possibilities, that which assigns the smaller value to I_a gives θ the smaller value, but neither assignment leads to values of α and μ , computed by means of the Debye formulas just given, that accord with the experimental values (see Table 16). Indeed, W. G. Penney and G. B. B. M. Sutherland 70 have concluded that attempts to obtain the exact form of the molecule from spectroscopic data are unprofitable. But numerous attempts of that kind have been made. From them it was at first concluded that I_a , and therefore θ , should have the smaller of the two possible values, this choice yielding values of

Table 16.—Constants of the Triangular Model of H2O

The atoms are at the vertices of an isosceles triangle, the O being at the unique vertex of angle 2θ ; a = altitude, b = base joining the two H's, r = slant height of the triangle. The principal moments of inertia are I_a about the axis parallel to a, I_b about that parallel to b, and I_c about that perpendicular to the plane of the triangle, all three axes passing through the center of mass of the system. For a rigid system $I_c = I_a + I_b$; for a nonrigid system, $I_o = I_a + I_b$ for the unvibrating molecule, and $I_o = I_a + I_b + I_b$ Δ for the vibrating molecule. The I's, being obtained from spectroscopic data, refer to the vibrating molecule and Δ is a positive quantity, generally differing from zero. In that case, Mecke and his associates derived θ from the values of I_a and I_b , but r from $I_1 + \Delta$ and I_2 , regarding $I_1 + \Delta$ as playing the part played by I_1 in the static molecule; here I_1 is the smaller and I_2 is the greater of the moments I_a and I_b . For the static molecule H_2O and cgs-units, $10^{-24}a^2 = 0.3364I_b$, $10^{-24}b^2 = 1.194I_a$, $10^{-24}r^2 = 0.3364I_b +$ $0.2985I_a$, $\tan \theta = 0.9416(I_a/I_b)^{0.5}$; for the associated triangle obtained by interchanging the values assigned to I_a and I_b , the elements are a' = b/1.884,

$$b' = 1.884a$$
, $r' = r \left[1.1269 - 0.2395 \left(\frac{a}{r} \right)^2 \right]_1^{0.5}$, $\tan \theta' = 0.8873 \cot \theta$. Until

recently, it was not obvious from the spectral data which of the two smaller moments should be assigned to I_b , but Mecke has concluded that I_b must be the smaller, which makes $2\theta > 90^\circ$. Previously, the other choice was preferred. The constants of both of the possible triangles are here tabulated; some of them have been computed by the compiler from the data given in the sources indicated.

⁷⁰ Penney, W. G., and Sutherland, G. B. B. M., Proc. Roy. Soc. (London) (A), 156, 654-678 (1936).

Table 16—(Continued)

The polarizability (α) and the dipole moment (μ) as computed from these values by means of Debye's formulas (see p. 51) $\alpha=(r^2/b)^3$, $\mu=2ea\left[1-(r/b)^3\right]$, do not agree with the observed values. For the obtuse-angled triangles 10^{24} α varies from 0.2 to 0.3 cm³ and 10^{18} μ from 3.9 to 4.4 cgse; for the acute-angled ones, 10^{24} α varies from 0.3 to 1 cm³, and 10^{18} μ from 0.7 to 3.2 cgse. The observed values are 10^{24} $\alpha=1.59$ cm³, 10^{18} $\mu=1.83$ cgse.

Unit	of r.	a, b	- 10-8	cm	-	1A
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			10C 611KIC			Acute angle			
Source*	θ	r	a	<i>b</i> '	66	r	a	ь	
FM	52.3°	0.952	0.583	1.507	34.5°	0.969	0.800	1.098	
FM'	52.5°	0.965	0.601	1.509	34.20	0.982	0.801	1.132	
LH	54.5	0.98	0.57	1.6	32.3	1.00	0.849	1.074	
VC	50	1.00	0.64	1.53	36.7	1.01	0.812	1,206	
EH	55.2	1.00	0.57	1.64	31.7	1.02	0.870	1.074	
ES	55	1.02	0.58	1.67	32	1.07	0.91	1.13	
W	56	0.99	0.55	1.63	30.9	1.02	0.865	1.036	
\mathbf{M}	48	0.86	0.58	1.28	38.6	0.87	0.680	1.093	
Pd	60								
Pl	57.5								

& Cources.

 α and μ that are the nearer to the experimental ones. But from an extended mathematical treatment J. C. Slater ⁷¹ has concluded that if the triangular configuration is to be in equilibrium, θ must slightly exceed 45°, which lies

EH = A. Eucken 18 and F. Hund 748; I = 0.98, I = 2.25, $I = 3.20 \cdot 10^{-40}$ g·cm².

ES = E. Eucken, 15 R. Sänger, and O. Steiger. 42

FM = K. Freundenberg and R. Mecke. Stationary state, $I_1 = 1.009$, $I_2 = 1.901$, $I_3 = 2.908$ 10^{-40} g·cm².

FM' = Same as FM except that it is for the vibrating state $\sigma = \pi = \delta = 0$, $I_1 = 0.996$, $I_2 = 1.908$, $I_3 = 2.981$, $I_3 - (I_1 + I_2) = 0.077$; θ is derived from $I_1 = 0.996$ and $I_2 = 1.908$; r, a, and b from $I_1' = 0.996 + 0.077 = 1.073$ and $I_2 = 1.908$.

LH = P. Lueg and K. Hedfeld.21

M = R. Mecke.78

Pd = J. Piccard.61

Pl = E. K. Plyler. 76

VC = J. H. van Vleck and P. C. Cross.77

W = Spectral data of H. Witt 78; $I_1 = 0.91$, $I_2 = 2.23$, $I_3 = 3.14$; the values of the I's and of the data in the table were computed by the compiler.

^b From observations on the spectrum of laterally scattered light, J. Cabannes and A. Rousset ⁷⁰ infer that 2θ [θ (?)] is about 23°, which does not agree with the other values.

[&]quot;Here the θ was computed from the $I_1 = 0.996$ and $I_2 = 1.908$, while a, b, and r were computed from $I_1' = I_1 + 0.077 = 1.073$ and $I_2 = 1.908$, which corresponds to another value of θ .

⁷¹ Slater, J. C., Phys. Rev. (2), 38, 1109-1144 (1931).

⁷¹ Wilson, E. B., Jr., J. Chem'l Phys., 4, 526-528 (1936); Randall, H. M., Dennison, D. M., Ginsburg, N., and Weber, L. R., Phys. Rev. (2), 52, 160-174 (1937).

⁷⁸ Mulliken, R. S., Phys. Rev. (2), 40, 55-62 (1932).

⁷⁴ Mulliken, R. S., J. Chem'l Phys., 1, 492-503 (1933); Idem, 3, 506-514, 586-591 (1935).

⁷⁴a Hund, F., Z. Physik, 32, 1-19 (1925).

⁷⁵ Mecke, R., Physik. Z., 30, 907-910 (1929).

⁷⁶ Plyler, E. K., Phys. Rev. (2), 38, 1784 (L) (1931); 39, 77-82 (1932).

⁷⁷ van Vleck, J. H., and Cross, P. C., J. Chem'l Phys., 1, 357-361 (1933).

⁷⁸ Witt, H., Z. Physik, 28, 249-255 (1924).

⁷⁰ Cabannes, J., and Rousset, A., Compt. rend., 194, 706-708 (1932).

between the two values fixed by the I's, and nearer the larger one. From spectral data, Mecke $^{13, 14}$ likewise concluded that only the larger value of the angle is satisfactory. Several sets of values that have been assigned to the elements of the triangular model are given in Table 16.

So far we have proceeded as if the molecule were rigid. It is not. Rotation causes distortion which is sufficient to produce marked spectroscopic effects under certain conditions.⁷²

The view now coming into favor is that of R. S. Mulliken,⁷⁸ who writes: "In general no attempt is made to treat the molecule as consisting of atoms or ions. Attempts to regard a molecule as consisting of specific atomic or ionic units held together by discrete numbers of bonding electrons or electron-pairs are considered as more or less meaningless, except as an approximation in special cases, or as a method of calculation. . . . A molecule is here regarded as a set of nuclei, around each of which is grouped an electron configuration closely similar to that of a free atom in an external field, except that the outer parts of the electron configurations surrounding each nucleus usually belong, in part, jointly to two or more nuclei..." The symmetry of the water molecule is that of an isosceles triangle, and the electron configuration, in spectroscopic notation, is given by him as $1s^22s^22pa^22pb^22pc^2$. "The order in which the symbols are written is that of decreasing firmness of binding." See also his article on electronic structure. The symmetry of the water molecule is that of decreasing firmness of binding. See also his article on electronic structure.

Association of the Molecules of Water-vapor.

Far from saturation, water-vapor behaves like an ideal gas with molecules of the composition H₂O.

As saturation is approached, both the density and the specific heat of water-vapor increase with abnormal rapidity, which indicates something of the nature of an association that increases as saturation is approached. The amount of association cannot be great, however, for the total departure of the density from that of an ideal gas is only a few per cent unless the pressure is high (see Section 14). The correct interpretation of these observations is difficult and not without a considerable degree of arbitrariness, as similar effects arise from the intermolecular forces that are taken into account by van der Waals' equation of state, though molecular aggregations caused by them are generally thought to be too transitory to be considered as associated molecules. Whether this distinction is justified is not entirely clear. As a result, interpretations differ, and it is necessary to consider in each case the assumptions on which the interpretation rests. In reference to the general subject, papers by J. W. Ellis ⁸⁰ and by E. J. M. Honigmann ⁸¹ are of interest.

It is generally assumed that if there is an association, the composition of the vapor is indicated by the expression $(1-x)H_2O + x(H_2O)_2$,

⁸⁰ Ellis, J. W., Phys. Rev. (2), 38, 693-698 (1931).

⁸¹ Honigmann, E. J. M., Dic Naturwiss., 20, 635-638 (1932).

x being the fraction of the molecules which are double. The ratio of the mass of double molecules to the total mass of vapor is 2x/(1+x).

From his observations at pressures not exceeding 80 per cent of that corresponding to saturation, and generally much lower, T. Shirai 82 concluded that x=0; he also concluded that the high values of x that had been computed by E. Bose 83 are not acceptable. The latter conclusion is reached also by A. W. C. Menzies. 84 Much earlier, S. Weber 85 had concluded that the molecular weight of water-vapor is 20 at -80 °C, and M. Knudsen 86 that it was 21.1 at -75 °C.

Assuming that the entire departure of the density of water-vapor from that of the ideal gas of molecular weight 18.0154 is due to the presence of double molecules, changing in number with the temperature and the pressure, H. Levy ⁸⁷ constructed an equation of state which satisfactorily represented the available data for the density and the specific heat; and W. Nernst ⁸⁸ derived in the same manner the extent of the association in the saturated vapor.

On the same assumption, but by a different procedure, A. Battelli ⁸⁹ had previously derived from his own observations corresponding, but markedly different, values, those for the higher temperatures being impossible.

- H. L. Callendar ⁹⁰ concluded that the density of superheated water-vapor can be expressed by an equation of state of the form $p(v-b)=RT-v_op$, in which $v_o=26.3(373/T)^{10/3}$ cm³/g. He calls v_o the "coaggregation volume," but does not interpret it more particularly; he takes b=1 cm³/g. If v_o is interpreted as the amount by which the specific volume is reduced by the formation of $(H_2O)_2$ from H_2O , then $x=v_o/(v-b)$. M. Jakob has discussed Callendar's ideas and theory in some detail, ⁹¹ and so has J. H. Awbery. ⁹²
- O. Maass and J. H. Mennie,⁹³ following a different procedure, found still other values.

Values of x corresponding to each of these several procedures are given in Table 17.

In a short note presented before the American Physical Society, H. T. Barnes and W. S. Vipond ⁹⁴ announced observations indicating that the

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82 Shirai, T., Bull. Chem. Soc., Japan, 2, 37-40 (1927).
83 Bose, E., Z. Elektrochem., 14, 269-271 (1908).
84 Menzies, A. W. C., J. Am. Chem. Soc., 43, 851-857 (1921).
85 Weber, S., Comm. Phys. Lab. Leiden, 150, 3-52 (1915).
86 Knudsen, M., Ann. d. Physik. (4), 44, 525-536 (1914).
87 Levy, H., Verh. deut. physik. Ges., 11, 328-335 (1909).
88 Nernst, W., Idem, 11, 313-327, 336-338 (1909).
89 Battelli, A., Ann. chim. phys. (7), 3, 408-431 (1894).
80 Callendar, H. L., "Properties of Steam," 1920.
81 Jakob, M., Engineering (London), 132, 143-146, 651-653, 684-686, 707-709 (1931).
82 Awbery, J. H., Rep. Prog. Phys. (Phys. Soc. London), 161-197 (1934).
83 Maass, O., and Mennie, J. H., Proc. Roy. Soc. (London) (A), 110, 198-232 (1926).
84 Barnes, H. T., and Vipond, W. S., Phys. Rev., 28, 453 (A) (1909). See also, Barnes, H. T.,
85 ("Ice Engineering," pp. 32-33, Montreal, Renouf Publ. Co., 1928.
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vapor arising from dry ice is initially polymerized to the same extent as ice itself, but quickly breaks down to water-vapor of the usual type, with the absorption of about 80 calories of heat per gram (335 joules/g).94 It seems that the details of this work have not been published. No other article suggesting anything of the kind has come to the compiler's attention.

From a study of the variation in the Raman spectrum with the density and temperature. S. A. Ukholin 95 has concluded that the molecules of water-vapor vibrate as if uninfluenced by their neighbors if the mean distance between them exceeds 10A (0.001 μ), corresponding to a density of 0.03 g/cm³. But if the mean distance does not exceed 8A ($\rho \equiv 0.06$ g/cm³) then there is an interaction; some of the molecules remain in the immediate vicinity of others for an interval that is long as compared with the period of the spectral vibration.

Table 17.—Estimates of the Extent of the Molecular Association of Saturated Water-vapor

(See text* for references and remarks)

The composition of the vapor is assumed to be $(1-x)H_2O +$ $x(H_2O)_2$, 2x/(1+x) and (1-x)/(1+x) being the fractions of the total mass that consist of double and of single molecules, respectively.

		Temperature - :	°C.		
Source→	Battellı 1894	Nernst 1909		Callendar	Maass and Mennie, 1926
,			-100x		
0	0.6	0.05		0.04	
10	0.7	0.08		0.06	
20	1.0	0.14		0.10	
50	1.2	0.41		0.35	
98				1.64	0.95
100	1.8	1.75		1.6	
108				1.40	0.71
200	7.7			9.5	
350	> 200			60.	

a Pressure is 1 atm.

10. Interaction of Water-vapor and Corpuscular Radiation

Alpha Particles.*

In water-vapor, the range of the alpha particles from polonium is 0.77 times their range in air at the same temperature and pressure.96 As their range in air at 0 °C and 1 atm is 3.72 cm 97 and varies inversely as the density, their range in water-vapor at T °K and a pressure of p mm-Hg is

^{*}For the very high estimates at -70 to -80 °C. by Knudsen and by Weber see p. 55.

^{*} Data from Kleeman, R. D., Int. Crit. Tables, 1, 370 (1926).

²⁶ Ukholin, S. A., Compt. rend. Acad. Sci. URSS, 16, 395-398 (1937). ²⁶ v. d. Marwe, C. W., Phil. Mag. (6), 45, 379-381 (1923). ²⁷ Geiger, H., Z. Physik, 8, 45-57 (1921).

R = 7.98T/p cm. Whence have been computed the following ranges (R) in water-vapor saturated at the temperature t:

*	0	15	20	25	50	100	°C
R^*	476	180	133	100	27.9	3.92	cm

^{*} More recent observations ** indicate that the range of these particles in air at 0 °C and 1 atm is 3.690 ± 0.005 cm; hence R for water-vapor is in each case 0.81 per cent smaller than the value here tabulated.

Electrons.†

When an electron strikes a molecule it may become attached to it, if the velocity of the electron is low, forming a negative ion; but if that velocity is great, the molecule is more or less disrupted, or ionized, and radiation may be emitted.

Ionization potential and energy.—The energy required to ionize a molecule is generally expressed in terms of the potential difference (I) through which an electron must pass in order to be able to cause the ionization. That difference I is called the ionizing, or ionization, potential. If I is expressed in volts, the energy required for the ionization is I electron-volts = $1.59 I \times 10^{-19}$ joule per ionized molecule = 96.4 I kj (= 23.0 I kcal) per gfw-H₂O.

When water-vapor is bombarded by electrons, ions of numerous types are formed, depending upon the energy of the electrons. As this was not at first recognized, and the nature of the ions formed was not determined, it is not surprising that some marked differences exist between the values of the ionization potential reported by the various early workers.

Finding that the transfer $A^+ + H_2O \rightarrow H_2O^+ + A$ occurs so readily that it is impossible to eliminate the ion H_2O^+ however carefully the argon is dried, H. D. Smyth and E. C. G. Stueckelberg ¹⁰⁴ concluded that either there is another ionization potential (well above 13 volts) corresponding to the removal of a different electron, or the collision in this case occasions an excitation to a higher level than that corresponding to ionization by electron impact. They were inclined to the second view, but at a later date Smyth ⁹⁹ preferred the first.

The energies required for removing the several individual electrons from H₂O have been estimated by R. S. Mulliken ¹⁰⁵ to be as follows, the electrons being designated in accordance with the notation currently used by spectroscopists: 2pc, 13.2 electron-volts (observed); 2pb, 2pa, and 2s, 16, 17 and 30 electron-volts, respectively; see also R. S. Mulliken.¹⁰⁶

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† See also Ionic Dissociation, Sections 7 and 8.

<sup>88</sup> Kurie, F N. D., Phys. Rev., (2), 41, 701-707 (1932).

<sup>80</sup> Smyth, H. D., Rev. Modern Phys., 3, 347-391 (384, 389) (1931).

<sup>80</sup> Mohler, F. L., Int. Crit. Tables, 6, 72 (1929).

<sup>81</sup> Mackay, C. A., Phys. Rev. (2), 24, 316-329 (1924); Phil. Mag. (6), 46, 828-835 (1923).

<sup>82</sup> Barton, H. A., and Bartlett, J. H., Jr., Phys. Rev. (2), 31, 822-826 → 154-155 (A) (1928).

<sup>83</sup> Smyth, H. D., Rev. Modern Phys., 3, 347-391 (385) (1931).

<sup>84</sup> Smyth, H. D., and Stueckelberg, E. C. G., Phys. Rev. (2), 32, 779-783 (1928).

<sup>85</sup> Mulliken, R. S., Phys. Rev. (2), 40, 55-62 (1932).
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Table 18.—Ionization Potential and Energy: Water-vapor

I = ionization potential, E = ionization energy, each for the particularion indicated; Int. = relative intensity of the indicated ionization when the pressure is about 0.5μ -Hg (= 0.0005 nm-Hg) and the speed of the electrons is that generated by a potential difference of 50 volts.

The data credited to S, MS, B, and L are those given by H. D. Smyth 99 as a result of his study of all the pertinent information available in June, 1931. SM states that the intensity of H₃O+ is approximately proportional to the square of the pressure; that of the other ions to the first power of the pressure.

Unit of I	= 1	volt; of	E =	10 kj/gfw-H ₂ O
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Ion	Int.	ı	E	Process	Ref.a
H ₂ O+		136	126	$H_2O \rightarrow H_2O^+$	S
H ₂ O+	1000	12.7 (13.3, 14.2, 15.0) 16.0	122-154		SM
OH+	200	18.9	182	$H_2O \rightarrow H + OH^+$	SM
OH+		17.3°	167		MS
H ₂ O+	200	Same as H ₂ O+	122-154	$H_2O^+ + H_2O \rightarrow H_3O^+ + OH$	SM
H ₃ O+		13.0	126		MS
H+	200	13.5, 18.9	130, 182	$H \rightarrow H^+; H_2O \rightarrow H^+ \rightarrow OH$	SM
H+		19.2	185		В
O+	20	18.5	178	$H_2O \rightarrow H_2 + O^+$	SM
O_2 +	6			$O_2 \rightarrow O_2^+$	SM
H ₂ +	6 5	33.5	323	$H_2O \rightarrow H_2^+ + O^+$	SM
OH-		d			SM
0-		e			SM
H-		6.6, 8.8 ^f	64, 85		L
Heat o	f forma	ation of H ₂ O	24		

Recent determinations of I

From spec	troscopic data	12.4	16.5			н
6.92	9.2	$ \begin{array}{r} 12.9 \\ 12.56 \pm 0.02 \end{array} $	16.7	17.8	24.5	R P
By electron	n impact	12.59 ± 0.05				SB
7.60			16.75			TW

a References:

- ferences:

 B Bleakney (reported by Loziei, q.v.).

 H Henning, H. J., Ann. d. Physik, (5), 13, 599-620 (1932).

 L Lozier, W. W., Phys. Rev. (2), 36, 1417-1418 (1930).

 MS Mueller, D. W., and Smyth, H. D., Idem, 38, 1920 (A) (1931).

 P Price, W. C., J. Chem'l Phys., 4, 147-153 (1936).

 R Rathenau, G., Z. Physik, 87, 32-56 (1933).

 S myth, H. D., Rev. Modern Phys., 3, 347-391 (384, 389) (1931).

 SB Smith, L. G., and Bleakney, W., Phys. Rev. (2), 49, 883 (A) (1936).

 SM Smyth, H. D., and Mueller, D. W., Idem, 43, 116-120 (1933) → Idem, 42, 902 (A) (1932).

 TW Thorley, N., and Whiddington, R., Proc. Leeds Phil. Lit. Soc. (Sci.), 3, 265-269 (1936). TW Thorley, N (1936).
- ^b F. L. Mohler 100 gives 13.2 volts on basis of observations by C. A. Mackay. 101
- . H. A. Barton and J. H. Bartlett, Jr., 102 reported 13 volts.
- d SM states that OH- first appears for I > 15 volts; as I increases, the intensity of OH-increases to a maximum, then decreases nearly to zero at 25 volts, then increases indefinitely.
- $^{\circ}$ SM states that as 1 increases, O- first appears at 22 ± 3 volts, increases to a maximum near 31 ± 4 volts, then decreases, to rise again at 36 ± 4 volts.
- 'Lozier could find II- only when I was very close to either 6.6 or 8.8 volts. "The significance of this remarkable result is not yet clear." SM found 7.9 volts for II-; they regard this as probably an unresolved combination of the two observed by L.

¹⁰⁶ Mulliken, R. S., J. Chem'l Phys., 3, 506-514 (1935).

¹⁰⁷ Gaviola, E., and Wood, R. W., Phil. Mag. (7), 6, 1191-1210 (1928).

E. Gaviola and R. W. Wood ¹⁰⁷ have concluded that about 5.2, and *not* less than 4.9, electron-volts are required to dissociate H₂O into H and OH. See also von Bishop, ¹⁰⁸ Grinfeld, ¹⁰⁹ and Townsend. ¹¹⁰

Ionization by accelerated electrons.—The number (α) of pairs of ions produced per cm of path by each electron of a stream driven through

Table 19.—Ionization of Water-vapor by Accelerated Electrons: Strong Fields

Adapted from the compilation by O. Stuhlman, Jr., ¹¹¹ from J. S. Townsend. ¹¹⁰

 α = number of pairs of ions produced '(= number of electrons freed) by each electron per cm of path in an applied uniform field of X volts/cm, the pressure of the vapor being p mm-Hg. Room temperature. Townsend derives the formula $(\alpha/p)_c = 12.9 \ e^{-289p/X}$.

X/p	100	200	300	400	500	600	700	800	900	1000
αp	1.31	3.6	5.2	6.35	7.2	7.95	8.5	9.0	9.4	9.7
$(\alpha/p)_c$	0.71	3.6 3.0	4.9	6.25	7.2	7.96	8.54	8.9	9.35	9.7

Table 20.—Miscellaneous Data for the Interaction of Accelerated Electrons with Water-vapor: Weak Fields

N. E. Bradbury and H. E. Tatel ¹¹² have reported that if p does not exceed 3 mm-Hg then no negative ions are formed if X/p is less than 10 (volt/cm) per mm-Hg, but that at higher values of p such ions are formed at lower values of X/p.

The following data are from V. A. Bailey and W. E. Duncanson.¹¹³ The electrons move in a uniform field of strength X; h is the probability of an electron's becoming attached to a molecule with which it collides; k is ratio of mean energy of agitation of the electrons to that of the molecules; L is mean free path of an electron when the pressure (p) of the vapor is 1 mm-Hg; u is mean velocity of agitation of the electrons; W is their mean energy, and w their mean velocity of drift; β is the probability of attachment in the interval of time required for the electron to move 1 cm along the direction of X; λ is the electron's fractional loss of energy during a collision with a molecule.

Unit of X=1 volt/cm; of p=1 mm-Hg; of $\beta=1$ in 100; of k=1 to 1; of k=1 and k=1 km/sec; of k=1 electron volt; of k=1 k=1 in 10000.

		electron. vc	nt; or L =	$I \mu$; or $\lambda =$	1%; OI n =	1 m 10000.		
X/p	β/⊅	k	าย	W	и	I.	λ	h
12	1.5	3.2	27	0.14	221	37	4.23	0.06
14	2.2	6.8	44	0.21	272	48	4.18	0.19
16	17	7.2	39	0.32	337	63	4.00	1.3
20	23	17	56	0.70	498	107	3.69	3.0
24	23	40	84	1.37	700	165	3.28	4.5
32	25	49	96	1.81	804	169	3.52	5.0

108 von Bishop, E. S., Physik. Z., 12, 1148-1157 (1911).
 109 Grinfeld, R., Univ. Nac. LaPlata, Estud. Cien. Fis. Mat., 4, 283-293 (No. 82) (1928); 4, 415-426 (No. 86) (1928); Physik. Z., 31, 247-252 (1930).

a gas by a uniform field (X) of sufficient strength to cause one electron to dislodge another from a molecule which it strikes varies with the density of the gas, but not otherwise with the temperature. If the pressure of the gas is p, then, at any given temperature, α/p depends solely upon X/p.¹¹¹ The factor by which the number of electrons in the stream is multiplied for each cm of its path is e^{α} , the number at the end of a path x cm long being $n = n_0 e^{\alpha x}$.

If the field is weak, the velocity of the impinging electron will be too low to dislodge another from the molecule, and it may itself be caught, forming a negative ion.

Excited Atoms and Molecules.

$$A^+ + H_2O \rightarrow H_2O^+ + A$$
, and $H_2O^+ + O_2 \rightarrow O_2^+ + H_2O^{-104}$

Collisions of excited Hg-atoms in the resonance level 2^3P_1 with normal water-vapor molecules may lead to several different processes. In most cases the Hg-atom is thrown down to the metastable $2^{3}P_{0}$ level, in a few cases (about 1 in 10000 collisions) the H₂O molecule is dissociated into H and OH, and in some cases (less than 1 in 1000) the complex quasimolecule Hg-H₂O is formed. The last dissociates, emitting a continuous band at 2800A.107 See also 114.

Mobility of Ions in Water-vapor.

The mobility (K) of an ion is its velocity of migration per unit field intensity. Over a wide range of densities, the product of K multiplied by the density is a constant for a given gas. The quantity K_0 satisfying the

formula
$$K_0 = K \frac{273p}{T}$$
, where K is the mobility observed at T °K and a

pressure of p atmospheres, has been called the "mobility constant" of the gas. The values found for K_0 are rather discordant, and opinions differ regarding the interpretation of the observed data. From recent data, it has been concluded that the mobility constant for normal ions in watervapor is $K_0 = 0.62$ cm·sec⁻¹ per volt·cm⁻¹ for the positive ion and 0.56 for the negative, but it is probable that each value should be increased by 20 per cent. 115 Numerical values are from L. B. Loeb and A. M. Cravath. 116

H. A. Erickson 117 finds that the molecule of H₂O in air gives up an electron to the final positive air ion, and thus forms an H₂O+ ion of greater mobility; and that the reciprocal of the mobility of negative ions in moist air is linear in the relative humidity.

¹¹⁰ Townsend, J. S., "Theory of Ionization of Gases by Collision," London, Constable, 1910. 111 Stuhlman, O., Jr., Int. Crit. Tables, 6, 121 (1929). From Bishop, E. S., Physik. Z., 12, 1148-1157 (1911).

¹¹² Bradbury, N. E., and Tatel, H. E., J. Chem'l Phys., 2, 835-839 (1934). 118 Bailey, V. A., and Duncanson, W. E., Phil. Mag. (7) 10, 145-160 (1930).

¹¹⁴ Senftleben, H., and Rehren, I., Z. Physik, 37, 529-538 (1926), and Bates, J. R., and Taylor, H. S., J. Am. Chem. Soc., 49, 2438-2456 (1927).

¹¹⁸ Loeb, L. B., Int. Crit. Tables, 6, 111 (1929).

¹¹⁸ Loeb, L. B., and Cravath, A. M., Phys. Rev. (2), 27, 811-812 (1926). 117 Erickson, H. A., Idem. 32, 792-794 (1928).

S. Chapman 118 has reported that when water is sprayed or when air is bubbled through water, ions of various mobilities are produced. When the numbers of ions are plotted against their mobilities one obtains as a background a broad flat curve with its maximum betwen the mobilities 0.05 and 0.10 cm/sec per (volt/cm), on which are superposed a number of In the sprayed liquid there are the same number of carriers of each sign, and the peaks occur at the mobilities 1.2, 0.3, and 0.2 for the negative carriers, and at 0.5 and 0.22 for the positive (first paper), or 1.5 for the negative and 0.9 for the positive (third paper). When air is bubbled through the liquid, the negative carriers are twice as numerous as the positive if the air tube is a small capillary, and 100 times as numerous if the tube is 12 mm in diameter (fourth paper); and the peaks occur at mobilities 1.2 and 0.25 for the negative and 0.7 and 0.3 for the positive (second paper), 1.5 and 0.3 for negative and 0.9 and 0.4 for positive (third paper), 1.9, 1.1, and 0.4 for negative and 1.1 and 0.4 for positive (fourth paper).

11. VISCOSITY OF WATER-VAPOR *

Four extended, but in part disagreeing, series of observations on the viscosity (η) of water-vapor have been reported: H. Speyerer (1925), range 1 to 10 kg*/cm², 107 to 347 °C; W. Schiller (1934), 1 to 30 kg*/cm², 100 to 300 °C; K. Sigwart (1936), 25 to 270 kg*/cm², 276 to 383 °C; and G. A. Hawkins, H. L. Solberg, and A. A. Potter (1935), 1 to 247 kg*/cm², 218 to 542 °C. Internal evidence indicates that the last is not satisfactory; for example, the effect of eddies in the wake of the falling body has been entirely ignored, although it must have been very perceptible in certain cases, and the instrumental temperature-coefficients used seem to be in error. It will not be considered further.

Of the first three, Speyerer and Schiller find that the values of η along an (η, p) -isotherm increase rapidly with the pressure (p), but the two sets differ markedly. Speyerer's values increase ever more rapidly as the pressure increases, whereas Schiller's exhibit marked irregularities. In contrast to them, Sigwart finds that η increases slowly and linearly with the pressure until the saturation pressure is rather closely approached, and then ever more rapidly. For the linear portion of the isotherms the slope is about 5 per cent per 100 kg*/cm², varying rather irregularly (2 to 7.6 per cent) from one isotherm to another. The total increase in η along the 275 °C isotherm to saturation does not exceed 1 per cent, and the increase to saturation is presumably less at lower temperatures. In striking contrast to this, Speyerer finds along the 270 °C isotherm an increase of 10 per cent on going from p = 1 to p = 10 kg*/cm², and Schiller finds 19 per cent for the same change.

¹¹⁸ Chapman, S., Phys. Rev. (2), 49, 206 (A) (1936); 51, 145 (A) (1937); 52, 184-190 (1937); 53, 211 (A) (1938).

^{*} For complete references see p. 68.

Under such conditions it is difficult to speak with confidence regarding the relative merits of the several series, but the author is inclined to favor Sigwart's. All three are given in Table 22, Schiller's values having been obtained by scaling his graph, which is a small one.

Values given by Sigwart for temperatures and pressures outside the domain covered by his observations were derived by extrapolation on the assumption that at 1 kg*/cm² $\eta = 16.47T^{1.5}/(T+548)$ micropoise (μp), that formula giving values to which his can be satisfactorily extrapolated. But that formula is based on H. Vogel's (1914) assumption that the Sutherland constant (the constant in the denominator) is to be taken as 1.47 times the absolute temperature of the normal boiling point, and that J. Puluj's (1878) reported value (90.4 μp) for η at 0 °C is correct. The first may be accepted; but the second should be 88.4, as Puluj used incorrect constants in reducing his observations (see Table 22, note b). Probably this difference of 2 per cent lies within the range of experimental error, but it should not be forgotten.

Fortunately, all three sets of values corresponding to 1 kg*/cm², observed or extrapolated, agree within \pm 2 per cent with one another and with the two short series by W. Schugajew (1934) and by C. H. Braune and R. Linke (1930), (see Table 23). So far as pressure is concerned, no distinction need be made between observations at 1 kg*/cm² and those at lower, but not excessively low, pressures.

No recent measurements at temperatures below 100 °C have been found; early ones are given in Table 22. Early observations were frequently made in terms of the viscosity of air, and were reduced to absolute units by assuming a value for air. In some cases the values were reduced to 0 °C on the basis of an assumed formula, generally requiring a knowledge of the absolute temperature (T_0 °K) of the ice point. Not infrequently an incorrect value for the viscosity of air or for T_0 was used, and sometimes an unsatisfactory formula. The resulting false values are commonly quoted, and appear in compilations of constants. An attempt has been made to correct such as are included in the following tables, the corrections that have been applied being explained in footnotes.

Formulas.

(a) Pressure = 1 kg*/cm^2 .—In the simple kinetic theory of gases the viscosity is given by the formula $\eta = BT^{1/2}$, where B is characteristic of the gas and is proportional to $nL_c = K/\pi d^2$, n being the number of molecules per unit of volume, L_c the effective mean free path of a molecule, d the molecular diameter (the diameter of the spheres by which the molecules may be regarded as replaced), and K a numerical factor determined by the distribution of the velocities of the molecules and by the kind of average free path L_c is. But actually η does not vary as $T^{1/2}$. Hence nL_c must vary with T. The failure of the formula may be explained by an attraction between the molecules, causing their paths to be curved, and increasing the frequency with which collisions occur. Since this curvature will decrease

as the translational kinetic energy of the molecules increases, Sutherland (1893) suggested that L_e should be replaced by $L_e/(1+CT^{-1})$, where C is a constant (Sutherland's constant), characteristic of the gas, which measures the intensity of the intermolecular attraction, and L_e is what the free path would be if there were no such attraction. Then $\dot{\eta}=AT^{1.5}/(T+C)$, A and C each being characteristic of the gas. This formula has been found to represent the observations on many gases over a wide range of temperatures. Similar expressions have been derived by S. Chapman (1916) and by J. H. Jeans (1916) with greater attention to mathematical rigor.

Unless observations are extended over a considerable range of temperature, C cannot be determined with precision. Consequently the experimental values that have been assigned to it vary greatly. Certain empirical formulas relating C to other quantities have been proposed. In 1910, A. O. Rankine observed that for the gases he investigated $T_{\rm crit}/C$ varied within a narrow range, the mean being 1.14; helium and hydrogen were marked exceptions. This relation suggested to Hans Vogel (1914) another between C and the normal boiling point (T_b) , and he concluded that $C = 1.47 T_b$. These and certain experimental values are given in Table 21.

Table 21.—Sutherland Constant for Water-vapor

$$\eta = AT^{1.5}/(T+C)$$

Observations by André Fortier (1936) indicate that for air the value of C depends on T.

C	Source
548 °K	H. Vogel (1914); $C = 1.47 T_b$
568	A. O. Rankine's relation (1910); C = T _{erri} /1.14
650	L. L. Bircumshaw and V. H. Stott, 123 from C. J. Smith (1924).
673	H. Speyerer (1925); experimental.
961	H. Braune and R. Linke (1930); experimental.

Actually, the observational data for water-vapor at and below $1 \, \mathrm{kg^*/cm^2}$ can, within experimental error, be represented essentially as well by means of a linear equation in the temperature (t °C) as by a Sutherland formula, and several observers give such equations. From a study of a great mass of data, Trautz (1931) concluded that $\eta = T^n \eta_{\mathrm{crit}}/T_{\mathrm{crit}}$, where n is, in general, a function of T, depends upon the type of substance, and approaches unity as T approaches T_{crit} . When n is unity, Trautz's formula is linear in t. An equation of this type with n=1 and $\eta_{\mathrm{crit}}=266 \, \mu \mathrm{p}$ has been accepted by some (see Sigwart) as fairly satisfactory for water-vapor at $1 \, \mathrm{kg^*/cm^2}$; and this in spite of the fact that it yields too large a value at $0 \, ^{\circ}\mathrm{C}$. With Sigwart's value for η_{crit} (378 $\mu \mathrm{p}$) a still higher zero value is obtained. The value $\eta_{\mathrm{crit}}=226 \, \mu \mathrm{p}$ (actually 228) seems to have been derived by Trautz in some unexplained way from the observations of others.

(b) Any pressure.—R. Plank (1933) has shown that Speyerer's data can be represented satisfactorily by the empirical formula (1) in which v* liters per gram is the specific volume of the vapor.

$$\eta = (86.1 + 0.373t) \cdot (1 + 0.0175 v^{*-1} + 0.0025 v^{*-2})$$
 micropoise (1)

Table 22.—Viscosity of Water-vapor

The values given in Section I are probably to be preferred in the domain t = 100 to 500 °C, P = 1 to 250 kg*/cm². For temperatures below 100 °C, see Sections V and VI. Sets of values for P = 1 are compared in Table 23.

Unit of $\eta=1$ micropoise (=1 μ p) = 1 μ g/cm·sec = 1.0197 × 10⁻⁸ kg*·sec/m²; of P=1 kg*/cm³ = 0.9678 atm. Temperature = t °C.

I. From K. Sigwart.¹²⁴ Values for P = 1 have been computed by means of the formula $\eta = 16.47T^{1.5}/(T + 548)$, which Sigwart regards as a satisfactory representation of the values obtained by extrapolating his observations. See text for bases of constants; T = 273 + t. Dubiety at P = 1 is 1 or 2 per cent; at 400 °C, 3 per cent; at 500 °C, 10 per cent; along saturation line, 1 per cent from 50 to 300 °C, 2 per cent at 360 °C, and 3 per cent at 370 °C. At critical point, $\eta = 378 \pm 5$ per cent.

$P \rightarrow t_{\text{sat}} \rightarrow t$	99.1	10 179.0	20 211.4	50 262.7	100 309.5	150 340 6	200 364.1	220 372.0	250
$t_{ m sat}$ 100	128 128 147	159	172	189	214	234	271	319	·
150 200 250	166 184	166 184	184						
300 350	201 219	202 220	203 221	206 225	231	238	0.00	070	201
400 450 500	235 252 268	236 238	237 239	238 241 241	244 243 243	248 245 245	256 249 248	272 257 251	286 263 255

II. From H. Speyerer. 125 He thinks that his observations at P=1can be satisfactorily represented by either of the formulas: $\eta = 88.33 +$ 0.3712t or $\eta = 86.8T^{1.5}/(T + 673)$; he uses the first.

$P \rightarrow t_{\text{mat}} \rightarrow$	99.1	119.6	142.9	158.1	8 169.6	10 179.0
t _{ent} 110	125.5 129.2	135.0	146.8	156.0	165.0	176.0
150 200	144.1 162.6	146.0 164.3	149.3 167.4	170.8	175.9	183.3
250 300	181.2 199.7	182.8 201.2	185.8 203.9	189.1 207.2	193.7 211.6	201.1 218.7
350	218.3	219.7	222.3	225.3	229.5	236.4

¹¹⁹ Mokrzycki, G., J. de phys. (6), 7, 188-192 (1926).

 ¹²⁰ Bircumshaw, L. L., and Stott, V. H., Int. Crit. Tables, 5, 6 (1929).
 121 Millikan, R. A., Phil. Mag. (6), 19, 209-228 (215) (1910).

¹⁹⁹ Stearns, J. C., Phys. Rev. (2), 27, 116 (A) (1926).

Table 22—(Continued)

III. From W. Schiller. 126 Values obtained by scaling his graph; he estimates the dubiety to be not over 2 per cent. At critical point, $\eta = 692$.

	$P \rightarrow$	Post	1	5	10	15	20	25	30
	Peak					77			
100	1.03	126	126						
120	2.02	141	133						
140	3.68	159	140						
150	4.85	170	144						
160	6.30	182	147	171					
180	10.12	208	155	174					
200	15.8	236	162	177	212	234			
220	23.6	268	170	183	215	235	255		
240	34.1	298	178	190	219	237	257	271	288
250	40.5		181	193	221	238	258	272	290
260	47.9		185	196	222	240	259	273	291
280	65.4		192	204	227	244	262	276	293
300	87.6		200	211	232	244	265	280	295
374	225.2	692							

IV. From W. Schugajew.¹²⁷ His mean of several discordant sets of observations. Within his limits of error, η does not vary with P over the range investigated (0 to 93).

t	100	150	200	250	300	350	400
77	126	144	164	183	202	222	241

V. From compilation by L. L. Bircumshaw and V. H. Stott¹²⁸; taken from a paper by C. J. Smith.¹²⁸ Various corrections have been made by the present compiler, and explained in the footnotes. Except as the contrary is indicated in the notes, the pressure was much less than 1 atm, and was far below saturation.

<i>t</i>	0 .	15	16.7	28.9	99.95	100	151.2	207.1
η	88.4 ^b	92.0°	94.7^{b}	99.7^{d}	125°	127	145	168
Rofe	D	KW	P	V	MS	Sm	Sm	Sm

VI. From H. Braune and R. Linke. ¹²⁹ These observations have not been published in detail, and the short table of only 11 lines of data referring to water-vapor contain at least two gross errors in computation, and one in transcription. These errors do not appear in the following values. Braune and Linke represent their observations by means of the formula $\eta = 22.36T^{1.5}/(T+961)$; the value (961) for the Sutherland constant is excessively high. At 20.2 °C the pressure was essentially $P_{\rm sat}$; for others, 120 to 210 mm-Hg.

t	20.2	92.6	107.5	210.3	313.1	366.4	406.6
77	93.7	117.8	124.2	163.8	214.9	226.1	242.2

¹²⁸ Bircumshaw, L. L., and Stott, V. H., Int. Crit. Tables, 5, 4 (1929).

¹²⁴ Sigwart, K., Forsch. Gebiete Ingenieurw., 7, 125-140 (1936).

¹²⁵ Speyerer, H., Forsch. Gebiete Ingenieurw., 273, 1-30 (1925) $\rightarrow Z$. Ver. Deuts. Ing., 69, 747-752 (1925).

¹²⁰ Schiller, W., Forsch. Gebiete Ingenieurw., 5, 71-74 (1934).

Table 22—(Continued)

- ^e References follow Table 23.
- $^{\circ}$ After reduction to the basis of $\eta = 179.2$ for air at 16.7 °C. ¹³⁰ The observations were made at 16.7 °C with an apparatus giving 183 for air at the same temperature, and yielded the value 96.7, which was published and is commonly quoted. The value 90.4 which he gives for water-vapor at 0 °C, and which is quoted in many tables, including the *Critical Tables*, was computed by him from his 96.7 value on the assumption that η is directly proportional to T and that 0 °C = 238 °K. On correcting the 96.7, using Sutherland's equation with C = 548, and placing 0 °C = 273.1 °K, one finds $\eta = 88.4$ at 0 °C.
- ° After reduction to the basis of $\eta=178.4$ for air at 15 °C. ¹³⁰ It was published and quoted as 97.5 at 15 °C, but on the basis of $\eta=189$ for air at 15 °C. The following entry in the *Critical Tables*, giving $\eta=97.5$ at 20.6 °C, was taken from Smith, who took it from Landolt-Börnstein, ¹³¹ where it had been doubly entered: once as the value at 15° C, and again as the value at 20.6 °C, the mean of the temperatures at which observations were taken. In the 5th edition of those Tabellen the value is entered but once, and is assigned incorrectly to 18.6-21.6 °C.
- ^d After reduction to the basis of $\eta=170.9$ for air at 0 °C. ¹⁸⁰ It was published and has been quoted in various tables, including the *Critical Tables*, as 100.6; but that was based on the assumption that $\eta=172.4$ for air at 0 °C. Pressure was near that of saturation.
- "After reduction to the basis of $\eta=170.9$ for air at 0 °C, as in note d. It was published and is commonly quoted as 132, but that is on the basis of $\eta=180$ for air at 0 °C. The pressure was near that of saturation.
- (c) Saturation pressure.—R. Plank (1933) also found that Speyerer's values for the viscosity (η'') of saturated water-vapor is related to that (η') of saturated water at the same temperature as shown by formula (2):

$$1/\eta'' + 1/\eta' = 1.1003 - 0.00261t \text{ micropoise}^{-1}$$
 (2)

But Schiller and Sigwart each find that this law of rectilinear mean is valid only over a limited range of temperatures.

Viscosity at the Critical Point.

Three estimates of the value of $\eta_{\rm crit}$ have been found: Trautz (1931) by an unknown method inferred that $\eta_{\rm crit} = 228.3 \,\mu{\rm p}$. By plotting and extrapolating the mean of the fluidities (η^{-1}) of saturated vapor and of saturated water at common temperatures, Schiller (1934) inferred 692, and Sigwart (1936) 378 $\mu{\rm p} \pm 5$ per cent.

Viscosity of Fog.

G. Mokrzycki ¹¹⁰ has concluded that the viscosity of foggy air is given by the relation $\eta = \eta_a + 1.59\Delta$ poise, where η_a (taken as 0.000171) is the viscosity of dry air, and Δ g/cm³ is the weight of the fog per cm³ of foggy air. In his observations, $10^6\Delta$ lay between 1.5 and 15, and the diameters of the fog particles between 0.5 and $10~\mu$. The value found for η was independent of the size of the particles.

¹²⁷ Schugajew, W., Physik. Z. Sowj., 5, 659-665 (1934).

¹²⁸ Smith, C. J., Proc. Roy. Soc. (London) (A), 106, 83-96 (1924).

Viscosity of Moist Air.

In their compilation,¹²⁰ L. L. Bircumshaw and V. H. Stott quote R. A. Millikan ¹²¹ as finding that air saturated at 26 °C has a viscosity 1904/1863 = 1.022 times as great as dry air at the same temperature, and J. C. Stearns ¹²² as claiming "that the viscosity of air is *decreased* by saturating it with moisture, the decrease being ½ per cent at 760 mm-Hg and 35 per cent at 14 mm-Hg pressures." It seems that this 6-line abstract is all that Stearns has ever published regarding his measurements. They are probably of little, if any, value.

Table 23.—Comparison of Various Values for the Viscosity of Water-vapor at or below 1 kg*/cm²

Several values from each of the sections of Table 22 are here compared with those defined by the formula $\eta_c = 16.12 T^{1.5}/(T+548)$, which gives Puluj's corrected value at 0 °C and uses Vogel's value for C.

The source of the data as well as the section of Table 22 in which they are given is indicated, and the individual observers of the several values given in *International Critical Tables* are designated. For additional information, see Table 22.

	1	Unit of $\eta = 1$ n	nicropoise.	t = temper	ature °C.		
Section of	rce→ f Table→	Sig.	Spe.	Schi.	Schu. IV	ICT V	BL VI
ı	ης				- ŋ		
0	88.4					88.4ª P	
15	94.1					92.0 KW	
16.7	94.7					94.7 P	
20.2	96.0						93.7
28.9	99.3					99.7 V	
92.6	123.0						117.8
99.95	125.8					125 MS	
100	125.8	128		126	126	127 Sm	
107.5	128.6	120		120	0	121 0111	124.2
150	144.1	147	144.1	144	144		1 ~ 2
151.2	144.5	177	177.1	111	177	145 Sm	
200	162.0	166	162.6	162	164	145 5111	
207.1	164.6	100	102.0	102	104	168 Sm	
						100 2111	162 0
210.3	165.7	404	101.0	101	102		163.8
250	179.6	184	181.2	181	183		
300	196.8	201	199.7	200	202		
313.1	201.9						214.9
350	213.6	219	218.3		222		
366.4	219.0						226.1
400	230.0	235			241		
406.6	232.1						242.2
500	261.7	268					

 a In the review 182 of the thesis of F. Houdaille (Paris, 1896) it is stated that he found 88.5 for water-vapor at 0 $^\circ$ C.

¹²⁹ Braune, H., and Linke, R., Z. physik. Chem. (A), 148, 195-215 (1930).

¹⁸⁰ Int. Crit. Tables, 5, 2 (1929).

¹⁸¹ Landolt-Börnstein, Phys.-Chem. Tabellen.

¹⁸⁸ Houdaille, F., Fortschr. d. Physik, 52, 442-443 (1897).

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KW Kundt, A., and Warburg, E., Ann. d. Physik (Pogg.), 155, 337-365, 525-550 (1875).
MS Meyer, L., and Schumann, O., Idem, (Wied.), 13, 1-19 (1881).
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Schi Schiller, W., Forsch. Gebiete Ingenieurw., 5, 71-74 (1934).
Schu Schugajew, W., Phys. Z. Sowj., 5, 659-665 (1934).

    Sig Sigwart, K., Forsch. Gebiete Ingenieurw., 7, 125-140 (1936).
    Sm Smith, C. J., Proc. Roy. Soc. London, (A), 106, 83-96 (1924).

Spe Speyerer, H., Forsch. Gebiete Ingenieurw., 273, 1-30 (1925) \rightarrow Z. d. Ver. deuts. Ing., 69, 747-752 (1925).
      Stearns, J. C., Phys. Rev. (2), 27, 116 (A) (1926).
      Sutherland, W., Phil. Mag. (5), 36, 507-531 (1893).
      Trautz, M., Ann. d. Physik (5), 11, 190-226 (1931).
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12. ACOUSTIC DATA FOR WATER-VAPOR

Velocity of Sound in Water-vapor.

In an ideal gas, the velocity of sound is independent of the pressure, and its square is directly proportional to the absolute temperature. For actual gases, especially near the region of liquefaction, departures from this relation should be expected. Nevertheless, many of the earlier data on the velocity of sound in gases and vapors were deduced on the assumption that this relation applies. Fortunately, it does apply fairly well for air at the temperatures that were used, and many of the observations consisted in determining merely the ratio of the velocity (v_t) in the gas or vapor under study at the temperature t to that $(u_{t'})$ in air at the temperature t'.

Then $\frac{v_t}{u_{t'}}\sqrt{\frac{T'}{T}}$ was called the ratio of the velocity at any temperature to that of air at the same temperature, and the product of this expression and the velocity in air at 0 °C was recorded as the velocity in the gas or vapor at 0 °C, although the observations may have been made at a quite different temperature. Actually, this ratio is not the ratio of the velocities at any temperature, but, so far as air satisfies the assumed relation, it is the ratio of the velocity in the gas or vapor at t to that in air at t. Hence the data can be reinterpreted when the value accepted by the observer for the absolute temperature of the ice-point is known; that can in certain cases be determined from the data he records. In some cases, $v_t \sqrt{T'}/u_{t'} \sqrt{T}$ has, incorrectly, been treated as if it were ratio of the velocity in the gas or vapor at t °C to that in air at 0 °C, thus leading to values that are entirely wrong.

The last seems to be the explanation of the excessively low values ($V_{98} =$ 402.4, $V_{96} = 410.0$ m/sec generally attributed to W. Jäger, and apparently originating in the Landolt-Börnstein Tabellen. They appear in the second and succeeding editions of the Tabellen (the first edition has not been examined) and in the International Critical Tables.

The first accounts for the value 401 m/sec given by A. Masson 188 for water vapor at 0 °C, and included in the tables mentioned. Actually, Masson's observations were made at 95 °C. Appropriate corrections have been applied to these in Table 24.

Table 24.—Velocity of Sound in Water-vapor

 $\delta_c = V_o - V$ and $\delta_i = V_i - V$, where V is the observed velocity, and V_{o} and V_{i} are those defined by the empirical formulas given in the text; T = 273.1 + t °C, the absolute temperature corresponding to t °C. Except as the opposite is indicated, the vapor was saturated at the temperature t °C.

	Ū	nit of V. &.	and 8; =	1 m/sec.	Temp. = 100	$T \circ K$		
	\boldsymbol{v}	V/\sqrt{T}	Obs.	*	\boldsymbol{v}	VIVT	8e	8,
27 100	432 405 ^b	24.94 21.0	Tha T		и	. G. Shillir	ngf	
Int.	Crit. Table	so (correcte	-	100	471.5	24.41	0.0	- 1.5
93.1	458.7 4	23.97	J	200	536.7	24.68	+ 1.2	+ 0.5
95 96.6	462° 467.7°	24.08 24.32	M T	300 400	593.2 643.2	24.78 24.79	+ 1.4 + 1.0	+ 0.7 0.0
100	471.5	24.41	Š	500	688.2	24.75	0.0	- 1.2
110 120	$\frac{413^b}{417.5^b}$	21.1 21.1	${f T}$	600 700	727.8 762.5	24.63 24.44	-0.1 + 0.8	-1.4 -0.1
130	424.4 ^b	21.1	$\dot{\mathbf{T}}$	800	795.3	24.28	+ 0.3	+0.3
1000	853.9	23.93	S	900 1000	825.0 853.9	24.08 23.93	-2.2	+1.4 + 1.3

- ^a G. E. Thompson. Frequency = 108.6 kilocycles/sec.
- ^b Velocity in tin tube, diameter = 1.4 cm. Pressure of vapor = 1 atm; hence the densities at 100, 110, 120, and 130 °C were, respectively, 1.00, 0.71, 0.51, and 0.38 of that corresponding to saturation at the indicated temperature.
- *From compilation by A. I., Foley ¹³⁰⁷ and ascribed, as indicated to: A. Masson, ¹³⁰⁸ W. Jäger, ¹³⁰⁹ W. Treitz, ¹⁴⁰ and W. G. Shilling, ¹⁴¹ For nature of corrections here applied, see preceding text.
- d These data of Jäger's are given in the Int. Crit. Tables and elsewhere as V∞ = 402, and $V_{10} = 410$; the densities of the vapor were, respectively, 0.55 and 0.70 of that corresponding to saturation at the temperature indicated.
- For this, most tables give Masson's $V_0 = 401$, though the observations were at 95 °C.
 - f Frequency = 3 kilocycles/sec.

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188 Masson, A., Ann. Chim. et Phys. (3), 53, 257-293 (1858).
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¹⁸⁴ Treitz, W., Diss., Bonn., 1903.

¹⁸⁴a Shilling, W. G., Phil. Maq. (7), 3, 273-301 (1927).

¹⁸⁵ Irons, E. J., Phil. Mag. (7), 3, 1274-1285 (1927).

 ¹⁸⁶ Thompson, G. E., Phys. Rev. (2), 36, 77-79 (1930).
 ¹⁸⁷ Foley, A. L., Int. Crit. Tables, 6, 461-467 (463) (1929).

¹⁸⁸ Masson, A., Compt. rend., 44, 464-467 (1857) → Phil. Mag. (4), 13, 533-536 (1857).

¹⁸⁹ Jäger, W., Ann. d. Physik (Wied.), 36, 165-213 (1889).

¹⁴⁰ Treitz, W., Diss. Bonn. (1903).

¹⁴¹ Shilling, W. G., Phil. Mag. (7), 3, 273-301 (1927).

The low values, $V_{110}=413$, $V_{120}=417.5$, and $V_{130}=424.4$ m/sec, attributed to W. Treitz ¹³⁴ in the tables mentioned are the velocities in tin tubes 14 mm. in diameter, and refer to superheated vapor at a pressure of 1 atm. His value $V_{100}=404.8$ m/sec for the saturated vapor in the same tubes seems to have escaped the attention of the compilers. It seems that he did not measure the velocity in air in those tubes.

The data of W. G. Shilling 134a were deduced by him from the velocity observed in a quartz tube 4 cm in diameter, the frequency being 3000 cycles/sec. They are approximately represented by the formula

$$V_o = 100 \sqrt{15.256 + 7.1135 \frac{t}{100} - 0.1384 \sqrt{\frac{t}{100}}}$$
 m/sec.

E. J. Irons ¹³⁵ represents the same data by the formula $V_i = -883.4 + 404.4T^{0.204}$ m/sec, where T = 273 + t °C.

Absorption of Sound by Water-vapor.

No information concerning the absorption of sound by pure water-vapor has come to the author's attention.

Moist Air.

The addition of water-vapor to air increases both the velocity and the absorption of sound. For any given frequency the absorption varies with the humidity, passing through a well-marked maximum; for any given humidity the absorption varies with the frequency, passing through a well-marked maximum at a certain frequency that varies as a quadratic function of the humidity. These maxima may be more than 20 times as great as the absorption to be expected from the classical theory. This effect is associated with the presence of oxygen; it does not appear with pure nitrogen. It is thought to arise from an acceleration, produced by intercollisions of O₂ and H₂O molecules, of the otherwise very sluggish process of equipartition of added energy among the several degrees of freedom of the oxygen molecule, a process which in the absence of H₂O requires a time of the order of 0.01 sec for completion. Collision with two H₂O molecules is very much more effective than collision with but one. (See ¹⁴²⁻¹⁴⁷.)

The following illustrative data for moist air at 20 °C, frequency 3000 cycles/sec, have been taken from the first of Knudsen's papers ¹⁴² just mentioned. Here w = number of H₂O molecules per 100 molecules of the mixture, and the absorption coefficient m is that defined for a plane wave

¹⁴² Knudsen, V. O., J. Acoust. Soc. Am., 5, 112-121 (1933) → Phys. Rev. (2), 43, 1051 (A) (1933).

¹⁴³ Kneser, H. O., Idem, 5, 122-23 (1933).

¹⁴⁴ Knudsen, V. O., Idem, 6, 199-204 (1935).

¹⁴⁵ Knudsen, V. O., and Obert, L., Idem, 47, 256 (A) (1935).

¹⁴⁶ Kneser, H. O., and Knudsen, V. O., Ann. d. Physik (5), 21, 682-696 (1935).

¹⁴⁷ Knudsen, V. O., and Obert, L., J. Acoust. Soc. Am., 7, 249-253 (1936).

by $I = I_0 e^{-mx}$ where x cm is the distance the wave travels while the intensity decreases from I_0 to I.

w 0.01	0.05	0.10	0.20	0.25	0.30	0.50	1.00	2.00
$10^4m \ 0.10$	0.32	0.74	1.82	1.95	1.68	0.82	0.47	0.33

The effect of moisture upon the velocity of sound in air is "comparatively slight."148 It was reported by C. D. Reid 150 to be represented at 20 °C by the formula $V_h = V_0 + 0.14h$, where h is the relative humidity; but that coefficient is in crror—it is ten times as great as his observations justify. He finds for dry air $V_0 = 331.68$ m/sec, for air saturated at 20 °C, $V_h = 333.05$, giving $V_h - V_0 = 1.37$ m/sec for h = 100; hence, at 20 °C, $V_h = V_0 + 0.0137h$. Since for saturation at 20 °C the partial pressure of H₂O is e = 17.51 mm-Hg, these values lead to $V_h = V_0$ (1 + Ae) where $10^4 A = 2.36$ per mm-Hg, the mean frequency being 130 kc/sec. This is not very different from the values found by others.

Table 25.—Velocity of Sound in Moist Air

The velocity (V_h) of sound in air in which the partial pressure of water-vapor is e mm-Hg may be conveniently represented by $V_h = V_d$ (1 + Ac), where V_d is the velocity in dry air, and A is an empirical coefficient that does not vary greatly with either the frequency, the temperature, or the humidity.

Unit of V and $\Delta = 1$ m/sec; of $\nu = 1$ kc/sec; of e = 1 mm-Hg; temp. = t °C

I. C. D. Reid. 160 At 20 °C, $\nu = 130$, $10^4 A = 2.36$ after correction (see text).

II. C. Ishii. 151 Observations at 30 °C.

ν	288	730	1439	2000	2892
10 ⁴ A	2.2_{3}	2.3	2.12	1.84	1.6

III. H. G. Muhammad.¹⁵² $\nu = 0.994$; air either dry (V_d) or saturated with H₂O (V_{sat}) ; $\Delta = V_{\text{sat}} - V_d$; values of A computed by compiler.

t	$V_{\mathtt{Sat}}$	17 a	Δ.	e	104A
15	341.55	340.40	1.15	12.78	2.64
20	344.83	343.51	1.32	17.51	2.18
25	348.10	346.35	1.75	23.69	2.13
30	351.5	349.3	2.2	31.71	2.0
35	355.3	352.1	3.2	42.02	2.2
40	259.10	354.95	4.15	55.13	2.12
45	363.05	357.75	5.30	71.6	2.07
50	367.2	360.7	6.5	92.3	1.95
55	371.7	363.4	8.3	117.8	1.93
60	376.9	366.2	10.7	149.2	1.96
65	382.8	368.9	13.9	187.3	2.01
70	389.8	371.6	18.2	233.5	2.10
75	398 6	374.3	24.3	299.1	2.16
80	408	377	31	355.1	2.3

¹⁴⁸ Hubbard, B. R., J. Acous. Soc. Am., 3, 111-125 (1931).

 ¹⁵⁰ Reid, C. D., Phys. Rev. (2), 35, 814-831 (1930).
 151 Ishii, C., Sci. Papers Inst. Phys. and Chem. Res. (Tokyo), 26, 201-207 (No. 560) (1935).

¹⁸⁹ Muhammad, H. G., Bull. Acad. Sci., Allahabad, 3, 269-294 (1934).

13. DIFFUSION OF WATER-VAPOR INTO GASES AND THROUGH SOLIDS

By definition, the coefficient of diffusion of a substance is the quantity D in expression (1)

$$\frac{dm}{d\tau} = -D\frac{d\rho}{dx}dA\tag{1}$$

in which dm is the mass of the substance that flows in the direction of increasing x through the area dA, perpendicular to x, in the time $d\tau$, when the gradient of the concentration of the substance (mass per unit of volume) is $d\rho/dx$. Expression (1) is equivalent to (2)

$$dm/d\tau = - (D \cdot d\rho/dp) \cdot dA \cdot (dp/dx) \cdot \tag{2}$$

in which p is the partial pressure of the substance at the point x.

If the gas is ideal, and for the present purposes water-vapor may be considered ideal, expression (2) can be given the following forms, in which R is the universal gas constant expressed in gram-moles and in the same units of p and x as are used in expression (2); A_n is the pressure of 1 atm, expressed in the same units as p; dv_0 is the volume of dm at 0 °C and 1 atm; $(p_m)_0$ is the pressure exerted by dm when at 0 °C and confined in a given volume V; and T_0 °K and T °K are the absolute temperatures at 0 °C and at the temperature at which the diffusion occurs.

$$dm/d\tau = - (DM/RT_0) \cdot (T_0/T) \cdot dA \cdot (dp/dx) \text{ g/sec}$$
 (3)

$$= - (D/RT_0) \cdot (T_0/T) \cdot dA \cdot (dp/dx) \text{ g-mole/sec}$$
 (4)

$$dv_0/d\tau = -\left(D/A_n\right) \cdot (T_0/T) \cdot dA \cdot (dp/dx) \tag{5}$$

$$(dp_m)_0/d\tau = -(D/V)\cdot (T_0/T)\cdot dA\cdot (dp/dx)$$
(6)

The units of mass in (3) and (4) are fixed by the specification that the unit of mass occurring in R shall be the g-mole. In the other equations the only restriction is that corresponding quantities shall be expressed in the same units and that the units of volume and of area shall be equal, respectively, to the cube and the square upon the unit of length.

For water-vapor the constant factors occurring in (3), (4) and (5) when the units of length, area, and volume are, respectively, 1 cm, 1 cm², and 1 cm³, and $T_0 = 273.1$, take the following values for each of the two units of p commonly used in such work.

Unit of	R	M/R	M/RT_0	$1/A_n$
1 atm	82.06	0.2195	8.039(10-4)	1
1 mm-Hg	62366	2.889(10-4)	$1.0577(10^{-6})$	$1.3158(10^{-8})$

Diffusion into Gases.

It has been found empirically that when one gas diffuses into another DP/T^n is constant, P being the total pressure and n being an empirically determined constant. Hence, if D_0 is the value of D at P_0 and T_0 , then the

value at P and T will be $D = D_0(T/T_0)^n(P_0/P)$. The value of n depends upon both the gases.

The subject, diffusion, has recently been discussed in a series of papers by M. Trautz and W. Müller, 153 with special reference to the corrections that must be applied to experimental data on account of various disturbing effects inherent in the procedures followed. They concluded that Winkelmann's data are the only ones capable of yielding reliable values for watervapor. See also H. Mache. 154

Table 26.—Diffusion of Water-vapor into Gases

The coefficient of diffusion (D) satisfies the following formulas: $dm/d\tau = -D(d\rho/dx)\cdot dA$; $DP = D_0P_0(T/T_0)^n$. Here dm is the mass crossing dA in time $d\tau$, P is the total pressure, and n is an empirically determined constant depending on both gases. See text for further information.

	Unit of	$D=1 \text{ cm}^2/\text{sec}$	Pressure $P=1$	atm. Temp. = t °C =	T °K
	Gas→		CO ₂	Air	H ₂
		and W. Mü	iller.155 Derived	from observations	by Winklemann.
See tex	ct.				
	$n \rightarrow$		2.115	1.853	1.8 44
	$D_0 \rightarrow$		0.1384	0.219	0.747
			± 0.0015	± 0.001	± 0.003
II.	Adapted fr	om compilati	on by W. P. H	Boynton and W. H	. Brattain. 156
	$n \rightarrow$	\boldsymbol{r}	2.00	1.75	1.75
	0	273	0.1387	0.220	0.7516
	10	283	0.1490	0.234	0.8004
	15	288	0.1544	0.242 6	0.8254

0	273	0.1387	0.220	0.7516
10	283	0.1490	0.234	0.8004
15	288	0.1544	0.242 b	0.8254
18	291	0.1576	0.246	0.8404
20	293	0.1598	0.249	0.8507
25	298	0.1653	0.257	0.8761
30	303	0.1708	0.264	0.9021
50	323	0.1939	0.295	1.009
70	343	0.2190	0.328	1.121
90	363	0.2452	0.362	1.238
100	373	0.2590	0.380	1.298

^a The values given by Boynton and Brattain are based upon the work of G. Guglielmo,¹⁵⁷ F. Houdaille,¹⁵⁸ M. LeBlanc and G. Wuppermann,¹⁵⁰ and A. Winkelmann.¹⁰⁰

^b W. E. Summerhays ¹⁶¹ has reported for air D = 0.282 at 16.1 °C and P = 1 atm.

Diffusion of Water-vapor through Solids.

In the case of a gas diffusing through a septum of thickness Δx and area dA, the total pressure being the same on both sides of the septum,

- 158 Trautz, M., and Müller, W., Ann. d. Physik (5), 22, 313-374 (1935).
- ¹⁵⁴ Mache, H., Sitzber. Akad. Wiss. Wien, (Math.-Nat. Abt. IIa), 119, 1399-1423 (1910).
- 185 Trautz, M., and Müller, W., Ann. d. Physik (5), 22, 333-374 (1935).
- 156 Boynton, W. P., and Brattain, W. H., Int. Crit. Tables, 5, 62-65 (1929).
- ¹⁸⁷ Guglielmo, G., Atti accad. sci. Torino, 17, 54-72 (1881); 18, 93-107 (1882) = Repert. d. Physik (Exner), 19, 568-581 (1883).
 - 158 Houdaille, F., Thesis, Paris (1896) → Fortschr. d. Physik, 52, 442-443 (1897).
 - 150 LeBlanc, M., and Wuppermann, G., Z. physik. Chem., 91, 143-154 (1916).
 - 100 Winkelmann, A., Ann. d. Physik (Wied), 22, 1-31, 152-161 (1884).
 - 161 Summerhays, W. E., Proc. Phys. Soc. London, 42, 218-225 (1930).

partial pressures of the gas in question being continually constant on each side and differing by Δp , the diffusion equation is that obtained by replacing dp/dx and dp/dx in formulas (1) to (5) by $\Delta p/\Delta x$ and $\Delta p/\Delta x$.

If the partial pressure of the gas in question is kept constant and equal to p_1 on one side of the septum, and increases on the other as a result of the accumulation of the transmitted gas in a vessel of fixed volume V, then the partial pressure (p_m) in that vessel will increase in accordance with expression (7)

$$dp_m/d\tau = (dp/d\rho) \cdot (d\rho/d\tau) = (dp/d\rho) \cdot (dm/d\tau)/V \tag{7}$$

If $dp_m/d\tau$ is small, and if the distribution of gas throughout the thickness of the septum is essentially the same as if the partial pressures on the two sides of the septum had for a long time been continuously the same as they actually are at the instant in question, then—and only then—may the $dm/d\tau$ in (7) be validly replaced by its value as given by (1), (2), (3), or (4), as modified in the way just stated. Using (2) and remembering that $-\Delta p = p_1 - p_m$. (7) becomes (7a)

$$dp_m/d\tau = (D \cdot d \cdot 1/V \cdot \Delta x) (p_1 - p_m) \tag{7a}$$

of which the solution is (8)

$$\log_{e} \left[(p_{1} - p_{0})/(p_{1} - p_{m}) \right] - D \cdot d \cdot 1 \cdot \tau / V \cdot \Delta x \tag{8}$$

in which p_0 is the value of p_m at $\tau = 0$.

Copper.—The data pertaining to the diffusion of water-vapor through copper and given in the compilation by F. Porter 162 are based on the observations by N. B. Pilling 163 and by H. G. Deming and B. C. Hendricks. 164 They consist of a value for D/D_h for Cu at 700 °C from the former, and $D_h T_0/T$ for Cu at 500 and at 750 °C from the latter, D_h being the value of the coefficient for H₂. The values that Pilling actually gives are the relative values of $dp_m/d\tau$ of expression (7) for each of several gases. The ratio of any two of those values will be equal to the ratio of the D's for the two gases only if the conditions were such as to justify expression (7a), if p_m were always negligible in comparison with p_1 , if p_1 were the same for each gas, and if the state of the copper were the same in both cases. The original article contains nothing regarding any of these items except the last. The same copper tube was used in all cases; but the purpose of the article was to prove that the structure of copper is profoundly modified by heating it in a reducing gas, and there is no indication of the order in which the observations on the diffusion of the several gases (H2, H2O, CO, CO₂) were made. Hence the value $D/D_h = 0.065$ at 700 °C inferred by Porter from those observations should be accepted only with great caution.

¹⁶² Porter, F., Int. Crit. Tables, 5, 76 (1929).

¹⁶⁸ Pilling, N. B., J. Franklin Inst., 186, 373-374 (1918).

¹⁰⁴ Deming, H. G., and Hendricks, B. C., J. Am. Chem. Soc., 45, 2857-2864 (1923).

The Deming and Hendricks paper contains a curve from which it is seen that H₂ passed through a certain septum of Cu at 700 °C at the rate of 0.020 mg/hr·cm² when $\Delta p/\Delta x = 1$ atm/nm; whence, by equation (3), $10^6D_h = 22$ cm²/sec at 700 °C. If this Cu septum was equivalent in structure to that used by Pilling, and if Porter's inference from Pilling's observations is justified, then for water-vapor and Cu at 700 °C, 10^6D = 22(0.065) = 1.43 cm²/sec, and $dm/d\tau = 320(10^{-12})$ g/sec·cm·atm.

Recently, J. H. deBoer and J. D. Fast 165 have reported for watervapor through Cu at 810 °C $dm/d\tau = 34(10^{-12})$ g/sec·cm·atm or 10^6D = 0.150 cm²/sec. This value is only slightly greater than 1/10 of that inferred in the preceding paragraph.

Table 27.—Permeability of Rubber to Water-vapor

(These sources are those quoted in Int. Crit. Tables, 2, 272, and 5, 76. See also Table 28.)

 $dm/d\tau = D \cdot dA \cdot (\Delta \rho/\Delta x), dv_0/d\tau = (DT_0/A_nT) \cdot dA \cdot (\Delta \rho/\Delta x), \kappa \equiv$ DT_0/A_nT , subscript h indicates that the quantity applies to hydrogen. For explanations of symbols, see text.

- J. Dewar 169 made some determinations on Para rubber. It was initially 1 mm thick, but was stretched until Δx was about 0.01 mm. One side of the rubber was in contact with water, which was considered as equivalent to the surface being in contact with saturated vapor. Under these conditions, at 15 °C, he found $\kappa/\kappa_{\rm air} = 163$. For another sample, also at 15 °C, he found $\kappa_{\text{nir}}/\kappa_h = 0.178$. Whence for water-vapor at 15 °C $\kappa/\kappa_h = 29.a$
- J. D. Edwards and S. F. Pickering 170 report data leading to the following values for a dental dam, for the two cases, assumed to be equivalent: (1) saturated vapor on one side; (2) water in contact with one side of the rubber.

	Unit of $\Delta r = 1$	mm, of D	- 1 cm ² /sec,	of $\kappa = 1$ cm	12/sec·atm. 1	°emp= 25 °C	
	(1) Va	por			(2) W	ater -	
Δx	$10^{5}D$	$10^6 \kappa^a$	K/A, b	Δx	10^6D	106Ka	κ/κ_h
0.18	23.5	21.5	47	0.21	46.4	42.5	95
0.25	28.1	25.7	62	0.25	51.9	47.2	115

^a In Porter's compilation ¹⁰² Dewar's value for κ/κ_h is given as 16, and the Edwards and Pickering values for 10% are given as 16.0 for (1) and 35 for (2). The values given in the table are those derived by the compiler from the data given in the original papers.

^b In Whitby's compilation ¹⁰⁸ the value of κ/κ_h for (1) is given as 55, essentially the mean of the two here tabulated.

deBoer, J. H., and Fast, J. D., Rec. Trav. Chim., Pays-Bas (4), 16, 970-974 (1935).

¹⁶⁶ Barrer, R. M., Nature, 140, 106-107 (L) (1937).

m Schumacher, E. E., and Feiguson, L., Ind. Eng. Chem., 21, 158-162 (1929).

¹⁰⁸ Whitby, G. S., Int. Crit. Tables, 2, 272 (1927).

Dewar, J., Proc. Roy. Inst. Grt. Brit., 21, 813-826 (1915).

170 Edwards, J. D., and Pickering, S. F., Sci. Papers Bur. Stand., 16, 327-362 (S387) (1920).

Chem. Met. Eng., 23, 17-21, 71-75 (1920).

Rubber.*—Certain remarks on the diffusion process in rubber have been published by R. M. Barrer. Since the publication of the International Critical Tables, E. E. Schumacher and L. Ferguson 167 have published their studies of the passage of water-vapor through rubber. The work was done under such conditions that expression (8) might be expected to apply. Their results are presented in the form of smooth curves of p_m versus τ . When values of p_m were read from those curves and log $(p_1 - p_m)$ was plotted against τ , the resulting graphs were not straight, but were convex toward the origin, showing that the observations do not satisfy expres-

Table 28.—Diffusion of Water-vapor through Various Solids

(For diffusion through copper see text; through rubber see also text and Table 27.)

 Δx = thickness of film; D = coefficient of diffusion; p = vapor pressure on "wet" side of film, on other side p = 0.

Unit of
$$D=1$$
 cm²/sec, of $p=1$ mm-Hg, of $\Delta x=1$ cm. Temp. = t °C

I. Acetylcellulose and nitrocellulose. 170a Not more than one film was used in a series; D varies with p, its value being determined by the amount of water in the film. The films were prepared on a polished plate (glass or silver), and stripped. The side that had been against the plate had a high polish; if that is the "wet" side of the film. D is greater than if the diffusion took place in the opposite direction; in a case for which data are given the difference was 14 per cent.

		lose, unfilled = 0.004 cm			
23.76 109	20.90 102	13.78 81	4.99 83	17.59 164	14.90 142
Acetylcellulose	filled with 1 $t = 25$ °C, Δ	part toluenesul $x = 0.0054$ cm,	famide to 1.5 $p = 23.756$ mi	parts ethylphtha n-Hg	late.
Filler (9 10 ⁶ D	%)	0 99	10 74	25 45	
Miscellane	ous mater	ials. ¹⁷¹			
Material		10 ⁸ ∆x	Δp	1	10°D
Asphalt sealing compound Balata Balata Bakelite (molded) Benzyl cellulose Cellulose acetate Cellulose acetate Cellulose film (waterproof) Phenol fiber Phenol fiber			22.8 10.5° 22.8 22.8 22.8 22.8 22.8 22.8 22.8 22.	25.0 25.0 25.0 23.9 25.0 25.0 25.0 25.0 25.0	3.30 5.16 4.82 13.8 30.3 448 465 3.39 234 14.0 15.0
	23.76 109 Acetylcellulose Filler (5 10°D) Miscellane Material sealing compo (molded) ellulose acetate acetate acetate acetate ber	$t = 25$ °C, $\Delta x = 6$ 23.76 20.90 109 102 Acetylcellulose filled with 1 $t = 25$ °C, Δ Filler (%) 10°D Miscellaneous mater Material sealing compound (molded) ellulose acetate acetate acetate (plasticized) efilm (waterproof) ber	109 102 81 Acetylcellulose filled with 1 part toluenesul $t = 25$ °C, $\Delta x = 0.0054$ cm, Filler (%) 0 99 Miscellaneous materials. 171 Material 10 ⁸ Δx sealing compound 82.0 38.4 (molded) 55.6 ellulose 68.6 acetate 16.3 acetate 15.6 acetate (plasticized) 2.45 efilm (waterproof) 4.40 ber 78.0	$I = 25 ^{\circ}\text{C}, \ \Delta x = 0.010 \text{cm}$ $23.76 \qquad 20.90 \qquad 13.78 \qquad 4.99$ $109 \qquad 102 \qquad 81 \qquad 83$ Acetylcellulose filled with 1 part toluenesulfamide to 1.5 $t = 25 ^{\circ}\text{C}, \ \Delta x = 0.0054 \text{cm}, \ p = 23.756 \text{m}$ Filler (%) 0 10 $10^{\circ}D \qquad 99 \qquad 74$ $Miscellaneous materials.^{171}$ Material 10* $\Delta x \qquad \Delta p$ sealing compound 82.0 22.8 $38.4 \qquad 22.8$ $38.4 \qquad 10.5^{\circ}$ (molded) 55.6 22.8 ellulose 68.6 22.8 ellulose 68.6 22.8 ellulose 68.6 22.8 elacetate 16.3 22.8 electate 15.6 22.8	$t = 25 {}^{\circ}\text{C}, \Delta x = 0.010 \text{cm}$ $t = 20 {}^{\circ}\text{C}, \Delta x = 20.000 \text{cm}$ $23.76 20.90 13.78 4.99 17.59$ $109 102 81 83 164$ Acetylcellulose filled with 1 part toluenesulfamide to 1.5 parts ethylphthat $t = 25 {}^{\circ}\text{C}, \Delta x = 0.0054 \text{cm}, p = 23.756 \text{mm-Hg}$ Filler (%) 0 10 25 $10^{\circ}D 99 74 45$ Miscellaneous materials. 171 Material 10^{\circ}\Delta \Delta \Delta p

^{*} See also Tables 27 and 28.

¹⁷⁰a Wosnessensky, S., and Dubnikow, L. M., Koll. Z., 74, 183-194 (1936).

¹⁷¹ Taylor, R. L., Herrmann, D. B., and Kemp, A. R., Ind. Eng. Chem., 28, 1255-1263 (1936).

Table	28(Continued)

Material	$10^3\Delta x$	Δρ		ŧ	10° D
Polystyrene $(10^3\Delta x \text{ from } 54 \text{ to } 209)$	Various	18.0	2	1.1	11.3
Polystyrene $(10^3\Delta x \text{ from } 54 \text{ to } 209)$	Various	11.90		1.1	11.2
Rubber hydrochloride (plasticized)	2.9	18.0		1.1	148
Vinyl chloride (plasticized)	48.1	18.0		1.1	10.7
Vinyl chloride (plasticized)	46.0	18.0	2	1.1	11.1
Wax, hydrocarbon	51	18.6	2	1.1	0.18
Rubber b (soft-vulcanized)	35.4	7.66		5.0	19.0
Rubber b (soft-vulcanized)	35.4	17.8		5.0	19.1
Rubber b (soft-vulcanized)	35.4	21.5	2.	5.0	19.7
Rubber ^b (soft-vulcanized)	35.4	22.8	25	5.0	21.0
Rubber b (soft-vulcanized)	35.4	23.6	2.	5.0	22.1
Rubber b (soft-vulcanized)	36.6	5.84	25	5.0	22.7
Rubber (hard; i. e., 32% combined					
Sulfur)	48.5	22.8	2.	5.0	4.33
Paragutta insulation ^c	46.9	22.8	2	5.0	5.51
Paragutta insulation ^c	44.9	22.8	2	5.0	5.34
Paragutta insulation ^c	46.7	10.5°	2	5.0	5.14
Paragutta insulation ^c	44.9	10.5	2	5.0	4.94
Guttapercha	32.4	22.8	25	5.0	4.25
Guttapercha	32.4	10.50	25	5.0	4.15
Chloroprene polymer (vulcanized)	86.5	18.0	2	1.1	7. 44
Polyethylene tetrasulfide	76.3	18.0	21	1.1	0.62
Temperature→ 0.0	21.0	21.1	25	30	35
Material		10° Z			
Polystyrene		11.3			13.3
Rubber (vulcanized) 13.2	19.5		21.0		25.2
Silk (varnished)				9.9	

^a The partial pressure of H_2O on one side of the material was always zero except for the cases to which this note refers; for them the partial pressure on the low-pressure side was as follows: for $\Delta p = 10.5$ it was 12.3; for $\Delta p = 11.9$, 6.1; for $\Delta p = 5.8$, 17.8.

sion (8). Furthermore, the values derived by the observers for what they call the permeability of rubber seem to have been based in each case on some kind of average of the slopes of the graphs over the first interval of 20 to 40 hours. Had an equal interval at a later time been used, much smaller values would have been obtained; their observations extended to 120 to 300 hours. For these reasons the reader is referred directly to their paper. They stated that D is inversely proportional to the thickness of the specimen.

For values reported in the *International Critical Tables* ^{162, 168} see Table 27.

Miscellaneous materials.—The articles cited in Table 28 contain much detailed data; they have not been critically examined by the compiler, who has merely converted the final values into the units here used.

^b Composition: Crepe 90, sulfur 1.5, zinc oxide 2.5, mineral rubber 3.0, paraffin 1.5, stearic acid 0.5, tetramethyl thiuram disulfide 0.5, phenyl- β -naphthylamine 1.0. Vulcanized in mold at 126 °C for 20 min.

Submarine cable insulation.

14. Pressure-volume-temperature Associations for Dilated Water-vapor

By dilated water-vapor is meant the vapor at a pressure that is less than its saturation pressure at the temperature concerned. The adjective "dilated" is less ambiguous than the more frequently used "unsaturated," which is equally appropriate to the supersaturated condition.

Equations of State.

Numerous equations of state for dilated water-vapor have been proposed, but none has been generally accepted (see ¹⁷²⁻¹⁸⁶).

Only the following equations of state are further considered in this compilation: Linde's (1905), equation (1), which represents very exactly the observations of O. Knoblauch, R. Linde, and H. Klebe, and in which $T_1 = 273.0 + t$.

Callendar's (1920-1928) equation (2), based upon quite different considerations, in which T = 273.1 + t,

$$Pv/m = 47T - P[0.0263(373.1/T)^{10/3} - 0.001] (kg/m^2) \cdot (m^3/kg)$$

= 4.551₂T - P[26.3(373.1/T)^{10/3} - 1] cm³·atm/g (2)

and Keyes, Smith, and Gerry's equation (3),188 representing their own observations at the Massachusetts Institute of Technology.

$$Pv/m = 4.55504T_2 + PB \text{ cm}^3 \cdot \text{atm/g}$$
 (3)

in which $T_2 = 273.16 + t$, temperature being t °C, and

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 184 Naumann, F., Z. physik. Chem. (A), 159, 135-144 (1932).
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 - 187 Knoblauch, O., Linde, R., and Klebe, H., Mitt. Forsch. Geb. Ing., 21, 33-55 (1905).

Table 29.—Isopiestics (kg*/cm²) of the Specific Volume of Dilated Water-vapor

(See also Tables 30 and 31).

The values given in the first section are those of Keyes, Smith, and Gerry, computed by means of formula (3) for $v/m > 10 \text{ cm}^3/\text{g}$, and derived by graphical methods for smaller volumes. In the second are the values obtained by J. Havliček and L. Miškovsky,¹⁹¹ and in the third are those derived by M. Jakob ¹⁸⁹ from the observed values of the specific heat at constant pressure. See also Table 32.

Unit of $p = 1 \text{ kg*/cm}^2 = 980665 \text{ dynes/cm}^2 = 0.96784 \text{ atm; of } v/m = 1 \text{ cm}^2/\text{g}$. Temp. = t °C

I. Keyes, S	Smith, and Ge	erry (1935)						
$p \rightarrow$	1	5	10		25		50	100
t				v/m —				
100	1729.6							
150	1975.4							
200	2215.8	433.8	210.4					
250	2454.1	484.1	237.6		88.99			
300	2691.3	533.2	263.3		101.1		46.41	
350	2927.9	581.6	288.2		112.1		53,12	23.03
400	3164.1	629.6	312.7		122.6		59.05	27.05
450	3400.1	677.4	337.0		132.7		64.60	30,41
500	3636.0	725.0	361 1		142.7		69.92	33.45
550	3871.8	772.5	385.1		152.6		75.10	36.32
$t \rightarrow t$	150	200	250		300		350	400
	11.98							
350	16.10	10.31	6.366		3.022^{a}		2.173ª	
400			9.456		6.979		5.168	3.858
450	18.90	13.05						
500	21.25	15.11	11.39		8.898		7.108	5.774
550	23.36	16.87	12.96		10.35		8.494	7.109
II. Havliček	and Miškov	rsky (1936)						
$t \rightarrow$	1	25		50		100		150
<i>t</i>	4520.5			v/m-				
100	1728.7							
150 200	1974.5 2215.1							
250 250	2453.4	88.96						
300	2690.5	101.0		46.38				
350	2927.2	112.0		53.07		23.02		11.97
400	3163.3	122.5		59.02		27.04		16.11
450	3399.4	132.7		64.58		30.41		18.91
500	3635.1	142.7		69.92		33.47		21.27
550	3870.9	152.6		75.10		36.34		23.40
$t \rightarrow$	200	250		300		350		400
t				-v/m				
400	10.31	6.365						
450	13.07	9.473		6.970				
500	15.13	11.42		8.927		7.126		5.761
550	16.91	13.01		10.40				

188 Keyes, F. G., Smith, L. B., and Gerry, H. T., Mech. Eng., 57, 164, 176 (1935); Proc. Am. Acad. Arts and Sci., 70, 319-364 (1935).

Table	20	(Continued)
Table	49-	Commune

III. Jakob	(1912)	2 4510 25	(Communa)		
p→	1	3	5	7	9
ŧ			v/m		
110	1781.6				
120	1830.2				
130	1878.9				
140	1927.3	630.5			
150	1975.5	647.6			
160	2023.7	664.6	392.3		
1 <i>7</i> 0	2 071.6	681.4	403.0	283.3	
180	2119.6	698.1	413.6	291.3	223.2
190	2167.4	714.6	423.9	299.2	229.6
200	2215 .2	731.1	434.2	306.8	235.9
220	2310.7	763.9	454.4	321.7	247.9
240	2406.0	796.4	474.4	336.4	259.7
260	2501.1	828.8	494.2	350.9	271.2
280	2596.0	861.1	514.0	365.3	282.6
300	2690.9	893.2	533. <i>7</i>	379.5	293.9
350	2927.9	973.3	582.4	414.7	321.7
400	3164.3	1052.9	630.6	449.6	349.1
450	3400.6	1132.3	678.6	484.2	376.2
500	3636.4	1211.3	726.2	518.4	402.9
550	3872.2	1290.2	773.8	552.5	429.5
$p \rightarrow$	11	13	15	17	19
<i>t</i>	1050		v/m		
190	185.2	150.1	1250		
200	190.6	159.1	135.9	1000	111.2
220	200.9	168.3	144.3	125.9	111.3
240	210.8	177.0	152.0	133.0	118.0
260	220.5	185.3	159.5	139.7	124.2
280	230.0	193.5	166.8	146.3	130.2
300	239.3	201.6	173.9	152.7	135.9
350	262.4	221.4	191.3	168.3	150.1
400	285.0	240.7	208.2 ^b	183.4	163.7
450	307.4	259.7	224.8	198.1	177.1
500	329.4	278.5	241.2	212.6	190.1
550	351.2	297.1	257.3	226.9	202.9

[•] It seems probable that one of these values given by Keyes, Smith, and Gerry for t = 400, p = 300, 350 is incorrect, see Table 32.

$$\begin{split} B &= B_0 + B_0{}^2g_1(\tau P) + B_0{}^4g_2(\tau P)^3 - B_0{}^{13}g_3(\tau P)^{12}, \ \tau = 1/T_2, \\ B_0 &= 1.89 - 2641.62\tau(10){}^{80870\tau^2}, \\ g_1 &= 82.546\tau - 1.6246\tau^2(10^5), \\ g_2 &= 0.21828 - 1.2697\tau^2(10^5), \\ g_3 &= 3.635(10^{-4}) - 6.768\tau^{24}(10^{64}). \end{split}$$

This equation and the data given in the Academy paper supersede all others previously published by these observers.

^b Published values 556.8 for p = 7, t = 550, and 210.9 for p = 15, t = 400, are obviously affected by typographical errors.

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¹⁹⁰a Wohl, A., Z. physik. Chem., 87, 1-39 (1914).

¹⁹¹ Havliček, J., and Miškovsky, L., Helv. Phys. Acta, 9, 161-207 (1936).

Table 30.—Isopiestics (lb*/in²) of the Specific Volume of Dilated Water-vapor 192

As published, the pressures refer to g = 981.16 cm/sec² (London); here, the unit so defined will be denoted by the subscript L; those without subscripts are based upon the international value, g = 980.665.

Unit of $p = 1(1b^*/in^2)L = 0.06898_2bar = 0.06808_0atm$; of	v/m = 1	$ft^a/lb = 62.428$	cm8/g.
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	Temp. = t °C								
	$t \rightarrow$	faat	250	300	350	400	450	500	
p	that				v/m -				
400	228.8	1.1712	1.2517	1.4269	1.5872	1.7391	1.8867	2.0292	
450	235.2	1.0419	1.0941	1.2562	1.4019	1.5392	1.6710	1.7991	
500	241.2	0.9380	0.9662	1.1191	1.2532	1.3790	1.4988	1.6151	
600	251.9	0.7805	0.7729	0.9124	1.0309	1.1387	1.2409	1.3392	
700	261.2	0.6672	0	0.7643	0.8709	0.9667	1.0560	1.1422	
800	269.7	0.5804		0.6515	0.7508	0.8379	0.9180	0.9943	
900	277.4	0.5120		0.5627	0.6569	0.7372	0.8102	0.8795	
1000	284.5	0.4570		0.4901	0.5816	0.6563	0.7239	0.7871	
1200	297.2	0.4370		0.4901	0.4673	0.5354	0.7239	0.6490	
				0.3707				0.5504	
1400	308.4	0.3111			0.3839	0.4482	0.5018		
1600	318.4	0.2635			0.3191	0.3823	0.4320	0.4762	
1800	327.5	0.2255			0.2661	0.3302	0.3775	0.4182	
2000	335.8	0.1936			0.2205	0.2879	0.3336	0.3719	
2400	350.6	0.1433				0.2216	0.2670	0.3022	
2800	361.3	0.1020				0.1703	0.2181	0.2517	
3200	373.6	0.0629				0.1253	0.1800	0.2136	
3600	0.0.0	0.000				0.0805	0.1491	0.1837	
4000						0.000	0.1213	0.1589	
1000							0.2410	0.2007	

Table 31.—Specific Volume of Dilated Water-vapor, and its Defect ¹⁹⁸ (Third International Steam-Table Conference, 1934.)

For the allowed tolerances and other data forming the skeleton tables then adopted for steam engineering, see Table 260.

 $\Delta \equiv 4.70636T/p - v/m$, T = 273.1 + t, units as stated below; $4.70636T \text{ (cm}^3/\text{g)} \cdot (\text{kg*/cm}^2) = 4.555T \text{ cm}^3 \cdot \text{atm/g}$. These values of Δ are comparable with those in Tables 32 and 33. Specific volume is v/m, pressure is p.

Unit of $p = 1 \text{ kg*/cm}^2 = 0.96784 \text{ atm}$; of $4.70636T = 1 \text{ (cm³/g)} \cdot \text{(kg*/cm}^2)$; of v/m and of $\Delta = 1 \text{ cm³/g}$. Temp. = t °C

			L CHIL/B. LCHI	p		
$p \rightarrow$	1	5	10	25	50	4 706267
<i>t</i>			v/m			4.70636T
100	1730					1755.9
150	1975					1991.3
200	2216	433.8	210.4			2226.6
250	2454	484.1	237.6	89.0		2461.9
300	2691	533.2	263.3	101.1	46.41	2697.2
350	2928	581.6	288.2	112.1	53.12	2932.5
400	3164	629.6	312.7	122.6	59.05	3167.8
450	3400	677.4	337.0	132.7	64.60	3403.2
500	3636	725.0	361.1	142,7	69.92	3638.5
550	3872	772.5	385.1	152.6	75.10	3873.8

¹⁹² Callendar, H. L., Proc. Inst. Mech. Eng., 1929, 507-527 (1929).

¹⁹⁸ Third International Steam-Table Conference, 1934; Mech. Eng., 57, 710-713 (1935).

					Table	31(Continue	ed)				
$t \rightarrow t$	_	75	10	00	125	5	150 v/m	20	00	250		300
300 350 400 450 500 550		27.48 33.22 37.78 41.83 45.62 49.25	27 30 33	.03 .05 .41 .45	16.6 20.5 23.5 26.1 28.5	3 2 4	11.98 16.10 18.90 21.25 23.36	13 13	0.31 3.05 5.11 5.87	6.36 9.46 11.39 12.96)	3.02 6.98 8.90 10.35
$t \rightarrow t$	1	5	10	25	50	75	100 - Δ	125	150	200	250	300
100 150 200 250 300 350 400 450 500 550	26 16 11 8 6 4 4 3 2	11.7 8.3 6.2 4.9 4.0 3.2 2.7 2.3	12.3 8.6 6.4 5.0 4.1 3.3 2.7 2.3	9.4 6.8 5.2 4.1 3.4 2.8 2.4	7.53 5.53 4.31 3.46 2.85 2.38	8.48 5.88 4.46 3.55 2.89 2.40	6.29 4.63 3.62 2.93 2.42	6.80 4.81 3.71 2.97 2.44	7.57 5.02 3.79 3.01 2.46	5.53 3.97 3.08 2.50	6.30 4.15 3.16 2.54	4.36 3.23

Table 32.—Defect of Specific Volume of Dilated Water-vapor: Isopiestics (kg*/cm²)

Here are assembled the values of the amount (Δ) by which the several values of the specific volume of water-vapor, as given in Table 29, fall short of that of an ideal gas (M=18.0154) under the same conditions of temperature and pressure. They may conveniently be used for computing the specific volume as defined by any of those sets of values and at any temperature and pressure within the range of the table.

 $v/m = 4.70636T/p - \Delta$, where T = 273.1 + t, units being as stated below; $4.70636T \text{ (cm}^3/\text{g)} \cdot \text{(kg*/cm}^2\text{)} = 4.555T\text{cm}^3 \cdot \text{atm/g}$, the basis used in this compilation.

Example: at 300 °C and 25 kg*/cm², $\Delta = 6.8$ for the K.S.G. set, and 6.9 for the H.M. set; that is, this specific volume in the second set is 0.1 cm³/g smaller than in the first, its actual value being (2697.21/25) - 6.9 = 101.0 cm³/g, agreeing with the corresponding H.M. value in Table 29.

Unit of $p = 1 \text{ kg*/cm}^2 = 980665 \text{ dynes/cm}^2 = 0.96784 \text{ atm; of } 4.70636T = 1 \text{ (cm³/g)-(kg*/cm²);}$ of $\Delta = 1 \text{ cm³/g}$. Temp. = t°(*

K	eyes, S	Smith,	and C	Gerry	(1935)							
•	1	5	10	25	50	100	150	200	250	30 0	350	400
	26.3						- 7					
	15.9											
	10.8	11.5	12.3									
	7.8	8.3	8.6	9.49								
	5.9	6.2	6.4	6.8	7.53							
	4.6	4.9	5.0	5.2	5.53	6.30	7.57					
	3.8	4.0	4.1	4.1	4.31	4.63	5.02	5.53	6.305	7.538°	6.878ª	
	3.1	3.2	3.3	3.4	3.46	3 62	3.79	3.97	4.157	4.365	4.555	4.650
	2.5	2.7	2.8	2.8	2.85	2.94	3.01	3.08	3.16	3.230	3.288	3.322
	2.0	2.3	2.3	2.4	2.38	2.42	2.46	2.50	2.54	2.56	2.574	2.576

Table 32—(Continued)

H	avliček and	d Mišk	ovsky	(1936)								
$p \rightarrow$	4 70C2CT	1	25	50	100	150) 	200	250	300	350	400
100 150 200 250 300 350 400 450 500 550	4.70636 <i>T</i> 1755.94 1991.26 2226.58 2461.90 2697.21 2932.53 3167.85 3403.17 3638.49 3873.80	27.2 16.8 11.4 8.5 6.7 5.3 4.5 3.8 3.4 2.9	9.52 6.9 5.3 4.2 3.4 2.8 2.4	7.56 5.58 4.34 3.48 2.85 2.38	6.30 4.64 3.62 2.91 2.40	7.5 5.0 3.7 2.99 2.4	8 1 8 9	5.53 3.95 3.06 2.46	6.306 4.140 3.13 2.48	4.374 3.201 2.51	3.268	3.335
Ja	ikob (1912).										
$t \rightarrow$		1	3	5		7	9	11	13	15	17	19
t	4.70636T	_						- Δ				
110	1803.00	21.4										
120	1850.07	19.9										
130	1897.13	18.2	?									
140	1944.20	16.9										
150	1991.26	15.8										
160	2038.32	14 6										
170	2085.39	13.8	3 13.			4.6						
180	2132.45	12.8		7 12		3.3	13.7					
190	2179.52	11.9			.0 1	2.2	12.6	12.				
200	2226.58	11.4				1.3	11.5	11.				
<i>22</i> 0	2320.71	10.0		7 9		98	100	10.				10.8
240	2414.83	8.8		5 8.		8.6	8.6	8.				9.1
260	2508.86	7.8				7.5	7.6	7.		7 7.8		7.8
280	2603 09	7.1				6.6	6.6	6.				6.8
300	2697.21	6.3		9 5.		58	5.8	5.				6.0
350	2932.53	4.0				4.2	4.1	4.				4.2
400	3167.85	3.6				3.0	2.9	3.				3.0
450	3403.17	2.0	2.			20	1.9	2.				2.0
500	3638.49	2 1				1.4	1.4	1.				1.4
550	3873.80	1.6	5 1.	1 1	.0	1.0^{b}	0.9	1.	0 0.	9 1.0	0 1.0	1.0

^a It seems probable that one of the values given for v/m at t = 400, p = 300, 350 (Table 29) is incorrect.

(Cont'd from p. 80)

Data computed by means of each of these equations will be found in the tables here given. In certain cases they are represented by the amounts by which they fall below the corresponding values defined by Pv/m = 4.555T cm³·atm/g. These defects, being strictly comparable, afford a ready means for comparing the several sets of values; they also facilitate interpolation.

By graphical means, M. Jakob ¹⁸⁰ has derived an extended table of specific volumes from the values found by Knoblauch and co-workers for the specific heat of water-vapor. ¹⁹⁰ These values agree well with the direct observations of Knoblauch, Linde, and Klebe, represented also by Linde's equation (1), and extend far beyond the range covered by them. They are given in Table 29.

Several sets of directly observed values are also tabulated, either directly

^b From the corrected values as given in Table 29.

or in terms of deviations. Values that have been accepted more or less by engineers at various times may be found in the numerous steam tables, see Table 259.

(To p. 90)

Table 33.—Defect of Specific Volume of Dilated Water-vapor: Isopiestics (atm)

Here are assembled the several values of the amount (Δ) by which certain observed and computed values of the specific volume of dilated water-vapor fall short of that of an ideal gas (M=18.0154) under the same conditions of temperature and pressure. They furnish a convenient means for comparing the several sets and for determining the specific volume, as defined by those sets, corresponding to any temperature and pressure within the range covered. At lower temperatures the relative departure from ideality, even at saturation, is small, $p\Delta/4.555T$ (= δ/v_c * of Table 250) being < 0.5 per cent if $t \ge 60$ °C, 0.9 per cent at 80 °C, 1.7 per cent at 100 °C, and 2.0 per cent at 110 °C.

Example: What is the value of pv/m at 160 °C and 3 atm on the basis of the "I" values (Int. Crit. Tables)? At 160 °C and 5 atm $\Delta = 15.2$, at 1 atm it is 15 - 1 = 14; whence at 3 atm it is 14.6, giving $p\Delta = 43.8$, and pv/m = 1972.8 - 43.8 = 1929.0. The value given in I.C.T. is 1929.

 $v/m = 4.555T/p - \Delta$ cm³/g; units as stated below.

Unit o	of $p=1$ at	m = 1.0	1325 ba	T = 1.0	3323 kg $3.1 + t$.	*/cm Te	mp. = t	$^{555}T = 1$	cm ³ atm/g	α , of Δ	1 cm³/g;
100∤→		17	21	30	37	45	57	61	67	68 7	2 73
t	4.555 <i>T</i>						Δ (Shira	ai) a			
80	1608.4		1		16						
100	1699.5	20		14		12			8 7		
120	1790.6			7						3.4	
140	1881.7								1	0 4	.3 5.0
$p \rightarrow$			1 ·							10	
Refa→		I	L						I	L	С
ŧ	4.555T						Δ				
110	1745.0	22	23	24							
120	1790.6	19	21	23							
130	1836.1	17	19	21							
140	1881.7	16		19							
150	1927.2	15	15	18							
160	1972.8		14	17	15.		15.8	15.4			
170	2018.3		12	16	13.	8	14.4	14.2			
180	2063.9		11	14			13.0	13.2			
185	2086.6		10	14			12.4	12.6	14.0	13.8	12.4
200	2155.0		9	13			10.8	11.2		11.9	11.0

^{*} References:

- C Callendar's equation of state, equation (2).
- I Compilation by F. G. Keyes 194 based on observations by A. Battelli, 195 M. Jakob, 189 and O. Knoblauch, R. Linde, and H. Klebe. 187
- L Linde's equation of state, equation (1).
- 5 T. Shirai. 196

¹⁹⁴ Keyes, F. G., Int. Crit. Tables, 3, 436 (1928).

¹⁹⁶ Battelli, A., Ann. Chim. Phys. (6), 26, 394-425 (1892); (7), 3, 408-431 (1894).

¹⁹⁶ Shirai, T., Bull. Chem. Soc. Japan, 2, 37-40 (1927).

Table 34.—Defect of Specific Volume of Dilated Water-vapor: Isopiestics (1b*/in²)

(From Callendar's data, see Table 30.)

 $v/m = 66.906_6 T/p - \Delta \text{ cm}^3/\text{g}$, the unit of p being $1 \text{ (lb*/in}^2)_L$ where $g = 981.16 \text{ cm/sec}^2 \text{ (London)}$; $66.906_6 T \text{ cm}^3 \cdot (\text{lb*/in}^2)_L/g = 4.555 T \text{ cm}^3 \cdot \text{atm/g}$. Hence these values of Δ are directly comparable with those of Tables 31, 32, and 33. For values of t_{sat} , see Table 30.

	Unit of	p = 1 (lb*/in	n²)L; of P	= 1 atm =	= 1.01325	bars; of A	= 1 cm ³ /g	
66.	t→ 906 ₆ T→	250 34998.8	300 38344,2	350 41689.5	400 45034.8	450 48380.1	500 51725.5	test
Þ	P				Δ-			
400	27.232	9.36	6.78	5.14	4.02	3.17	2.63	10.84
450	30.636	9.47	6.79	5.12	3.99	3.19	2.63	10.53
500	34.040	9.68	6.82	5.14	3.98	3.19	2.62	10.26
600	40.848	10.08	6.95	5.13	3.97	3.17	2.60	9.82
<i>7</i> 00	47.656		7.06	5.20	3.99	3.19	2.59	9.42
800	54.464		7.26	5.24	3.99	3.17	2.58	9.16
900	61.272		7.48	5.31	4.02	3.18	2.57	8.96
1000	68.080		7.75	5.38	4.06	3.19	2.59	8.78
1200	81.696		8.43	5.57	4.10	3.21	2.59	8.53
1400	95.312			5.81	4.19	3.23	2.59	8.37
1600	108.928			6.14	4.28	3.27	2 .60	8.28
1800	122.544			6.55	4.40	3.31	2.63	8.25
2000	136.160			7.08	4.54	3.64	2.65	8.28
2400	163.392				4.93	3.49	2.69	8.44
2800	190.624				4.45	3.66	2.77	8.83
3200	217.856				6.25	3.88	2.83	9.59
3600	245.088				7.48	4.13	2.90	
4000	272.320					4.52	3.01	

Table 35.—Isometrics of the Pressure (atm) of Dilated Water-vapor 197

Four sets of experimental data, here indicated by superscripts a, b, c, and d, as finally corrected by the authors, are given. These include, with corrections, and supersede similar data previously reported from that laboratory, ¹⁹⁸ and are the data from which formula (3) was derived.

Unit of	$v/m=1~\rm cm^3/g,$	of $P=1$	atm = 1.01325	bars = 1.03323	kg*/cm2.	Temp. = # °C
$v/m \rightarrow$	150.04	140.0b	100.00	97.5ª	75.00	75.00
t				– P ––––		
195		13.787				
196		13.898				
197		13.957				
198		14.003				
199		14.040				
200	13.220	14.074				
210	13.591	14,479				
212.5			19.612			
220	13.952	14.871	20.086	20.514		(26.238)
230	14.302	15.252	20.664	21.106	26.50	

Keyes, F. G., Smith, L. B., and Gerry, H. T., Proc. Am. Acad. Arts Sci., 70, 319-364 (1935).
 Smith, L. B., and Keyes, F. G., Mech. Eng., 52, 123-124 (1930); 53, 135-137 (1931); 54, 123-124 (1932); Keyes, F. G., and Smith, L. B., Idem, 53, 132-135 (1931).

		Tabl	e 35—(Cont	inued)		
$v/m \rightarrow$	150.04	140.0 ^b	100.08	97.5ª	75.0°	75.0b
240 250 260 270 280	14.647 14.986 15.323 15.658 15.990	15.627 16.014 16.366 16.745 17.087	21.217 21.769 22.297 22.825 23.356	21.688 22.258 22.816 23.365 23.905	27.309 28.099 28.869 29.618 30.355	27.303 28.082 28.854 29.592 30.345
290 300 310 320 330	16.319 16.646 16.971 17.297 17.620	17.448 17.802 18.146 18.502 18.839	23.863 24.378 24.881 25.391 25.892	24.437 24.964 25.485 26.001 26.514	31.081 31.794 32.496 33.188 33.870	31.055 31.767 32.476 33.154 33.857
$v/m \rightarrow$	57.54	50.08	50.0€	40.0	40.00	40.04
250 260 264 265	34.983 36.051	39.052 40.469	39.388 40.592	48.689	48.827 48.996	
270	37.090			49.753	49.855	49.856
280	38.104	42.988	43.088	51.417	51.511	51.533 53.123
290 300 310 320	39.101 40.078	45.366	45.483	54.613	54.708	54 665 56 169
	41.035 41.980	47.688	47.751	57.043	57.711	57.642
330 340 360 380 400	42.915	49.914 52.105 54.257 56.384	49.985 52.188 54.308 56.426	60.586 63.428 66.225 68.963	60.646 63.497 66.279 69.015	59.096 See end of table
420 440 460		58.465 60.546 62.605	58.503 60.555 62.608	71.658 74.311 76.966	71.682 74.320 76.947	
$v/m \rightarrow t$	39.56	30.0	30.0◦	20 0d	20.0	20.0 °
264 270 280	49.164 50.322 52.008			- F		
281 290	53.635	63.622 65.821	63.730 65.921			
300	55.229	68.145	68.220			20.404
305 305.5 310 320	56.745 58.261	72.568	72.611	90.765 92.582 96.342	90.499 96.449	90.604 (91.143) ' 96.301
330	59.718	72.306	72.011	100.009	90.449	90.301
340 350	010	76.832	76.841	103.586 107.103	103.627	103.654
360 380		80.841 84.804	80.900 84.845	110.525 117.105	110.456 117.049	110.504 117.092
400 420 440 460		88.674 92.437 96.161 99.836	88.678 92.439 96.152 99. 789	123.535 129.632 135.594 141.484	123.400 129.628 135.656 141.580	123.436 129.624 135.674 141.587

Table 35-	-(Continued)
Table 33	-(Committee)

$v/m \rightarrow$	17.5¢	15.0¢	12.54	10d	7.5ª	6.254
t				- P		
313	101.287					
320	104.286					
322	400 400	113.965				
330	108.690	118.413	100 500			
332.5			130.593			
340	112.988	123.725	135.657			
344.5				151.833		
350	117.154	128.838	142.215	156.607	450 400	
357.5					178.402	
360	121.235	133.857	148.566	165.1 <i>37</i>	181.389	
364.5						194.118
370			154.741	173.384	193.226	201.951
380	129.679	143.453	160.516	181.385	204.683	216.133
400	136.6 66	152.670	172.435	196.909	226.821	243.518
420	143.930	161.501	183.610	211.775	248.049	269.746
440	150.958	170.094	194.464	226.179	268.715	295.539
460	157.886	178.473	204.988	240.171	288.700	320.444
$v/m \rightarrow$	5 0d	4 0ª	3.0 ^d	2.0^{d}	$v/m \rightarrow t$	40 ^b
* 370				220.025	310	56,192
371.5	210.784			220.025	320	57.638
371.5 375	210.704	219.842	220.644		330	59.102
373 380	225.819	230.931	234.370	267,878	330	37.102
390	225.017	252.748	262.404	317.298		
400	260.715	274.367	290.947	367.654		
410	200.713	295.536°	319.801	307.034		
420	294,563	317.218	348.962			
420 430	234.303	338.537	340.902			
440	327.836	359.710				
460	360.173	007.710				
100	000.170					

a, b, c, d These superscripts serve to identify the several sets of data.

Table 36.—Isometrics of the Pressure $(kg*/cm^2)$ of Dilated Water-vapor 199

(See also Table 38.)

Experimental results as smoothed by the observers.

	Unit of	v/m = 1 cm	18/g; of p =	1 kg*/cm2	= 0.96784	atm. Temp	$o. = t ^{\circ}C$	
$v/m \rightarrow$	10	9	8	7	6	5.5	5	4.5
<i>t</i>	-				p			
350	162							
360	174	181						
370	184	193	202	209	213			
380	192	201	211	221	229	231	233	235
390	199	210	221	233	244	248	252	255
400	207	217	230	245	258	264	269	274

¹⁹⁹ Nieuwenburg, C. J., and Blumendal, H. B., Rec. trav. chim. Pays-bas, 51, 707-714 (1932).

[°] This is for t = 227.5 °C.

^{&#}x27;This is for t = 306.0 °C.

[&]quot;The authors state that this value is evidently in error; by interpolation they find 295.866.

Table 36—(Continued)											
10	9	8	7	6	5.5	5	4.5				
213 219 226	225 232 240 248	238 247 257 266	255 266 277	272 286 298	279 294 308 323	286 303 319 334	293 312 330 347				
239 246 252 258	254 262 269 277	275 284 292 297	298 310 319 328	323 337 347 359	337 351 363 376	351 366 380 395	364 382 399 416				
4	3.5	3.0	2.5	2.2	2.0	1.8	1.6				
236 259 281 301 322	238 262 287 311 334	241 268 296 324 351	246 280 313 348 381	256 300 339 379 423	280 329 379 430 477	344 403 462 520 582	511 600				
342 361 381 400 419 439	356 379 401 424 444 467	378 405 432 459 487 515	416 450 486 520 553 589	465 504 545 591	525 571						
	213 219 226 233 239 246 252 258 4 236 259 281 301 322 342 361 381 400 419	213 225 219 232 226 240 233 248 239 254 246 262 252 269 258 277 4 3.5 236 238 259 262 281 287 301 311 322 334 342 356 361 379 381 401 400 424 419 444	10 9 8 213 225 238 219 232 247 226 240 257 233 248 266 239 254 275 246 262 284 252 269 292 258 277 297 4 3.5 3.0 236 238 241 259 262 268 281 287 296 301 311 324 322 334 351 342 356 378 361 379 405 381 401 432 400 424 459 419 444 487	10 9 8 7 213 225 238 255 219 232 247 266 226 240 257 277 233 248 266 288 239 254 275 298 246 262 284 310 252 269 292 319 258 277 297 328 4 3.5 3.0 2.5 236 238 241 246 259 262 268 280 281 287 296 313 301 311 324 348 322 334 351 381 342 356 378 416 361 379 405 450 381 401 432 486 400 424 459 520 419 444 487 553	10 9 8 7 6 213 225 238 255 272 219 232 247 266 286 226 240 257 277 298 233 248 266 288 311 239 254 275 298 323 246 262 284 310 337 252 269 292 319 347 258 277 297 328 359 4 3.5 3.0 2.5 2.2 236 238 241 246 256 259 262 268 280 300 281 287 296 313 339 301 311 324 348 379 322 334 351 381 423 342 356 378 416 465 361 379 405	10 9 8 7 6 5.5 213 225 238 255 272 279 219 232 247 266 286 294 226 240 257 277 298 308 233 248 266 288 311 323 239 254 275 298 323 337 246 262 284 310 337 351 252 269 292 319 347 363 258 277 297 328 359 376 4 3.5 3.0 2.5 2.2 2.0 236 238 241 246 256 280 259 262 268 280 300 329 281 287 296 313 339 379 301 311 324 348 379 430 322<	10 9 8 7 6 5.5 5 213 225 238 255 272 279 286 219 232 247 266 286 294 303 226 240 257 277 298 308 319 233 248 266 288 311 323 334 239 254 275 298 323 337 351 366 252 269 292 319 347 363 380 258 277 297 328 359 376 395 4 3.5 3.0 2.5 2.2 2.0 1.8 236 238 241 246 256 280 344 259 262 268 280 300 329 403 281 287 296 313 339 379 462 301 31				

Table 37.—Defect of Pressure (atm) of Dilated Water-vapor: Isometrics

At the end of the table are certain values derived from the L (Linde) data of Table 33; the rest have been derived from the Keyes, Smith, and Gerry data of Table 35, the b data only being used when there are several sets referring to the same value of v/m.

The fractional defect is δ , defined by the relation $P = P_i(1 - \delta)$, where P_i (=4.555Tm/v atm) is the pressure of an ideal gas (M = 18.0154) at the same temperature and density, except for errors in the assumed value of the gas constant (R) and in T. Here T = 273.1 + t.

The values tabulated are $\delta(v^*+1)$, it having been observed that this product varies less with v/m ($\equiv v^*$) than does either δv^* or $\delta(v^*+2)$. These values enable one to compare readily this set of data with those from which the values in Table 38 were derived, and to derive the approximate value of P corresponding to any specific volume and temperature within the range of the table. For rough estimates of P, the proper value of $\delta(v^*+1)$ may be derived from the table by inspection; for more precise values, use may be made of the fact that $\delta(v^*+1)$ varies almost linearly with the density, t being constant; and almost linearly with 1/T, v/m being constant.

As δ has been derived directly from experimental data, it is subject to experimental irregularities, and so are the values of P derived from it.

Table 37—(Continued)

Unit of	$v/m \equiv v^* = 0$	1 cm8/g; of 4	1.555T = 1	cm8.atm/g;	δ is dimens	ionless. Te	mp. ='t °C
$v/m \equiv v^* \rightarrow$		150	140	100	97.5	75	57.5
# 195 196 197 198 199	4.555 <i>T</i> 2132.20 2136.75 2141.30 2145.86 2150.42		13.359 12.605 12.335 12.185 12.118	δ(υ*	+ 1)		
200 210 220 230 240	2154.97 2200.52 2246.07 2291.62 2337.17	12.050 11.107 10.304 9.642 9.053	12.078 11.115 10.304 9.619 9.013	10.679 9.926 9.311	10.786 10.053 9.381	10.131 9.412	
250 260 270 280 290	2382.72 2428.27 2473.82 2519.37 2564.92	8.544 8.073 7.637 7.245 6.893	8.329 7.957 7.240 7.118 6.717	8.724 8.259 7.811 7.366 7.034	8.787 8.263 7.793 7.375 7.001	8.821 8.270 7.817 7.345 6.987	9.114 8.560 8.067 7.625 7.221
300 310 320 330	2610.47 2656.02 2701.57 2747.12	6.569 6.275 5.982 5.723	6.384 6.136 5.809 5.629	6.670 6.385 6.074 5.806	6.659 6.350 6.070 5.808	6.636 6.304 6.049 5.750	6.857 6.531 6.230 5.952
$\begin{array}{c} t_{\mathtt{sat}} \rightarrow \\ P_{\mathtt{i}} \rightarrow \\ \delta(v^{+} + \\ \end{array}$	-1)→	191.90 14.210 12.913	195.26 15.238 12.542	211.98 22.095 11.466	213.33 22.725 11.304	227.30 30.391 10.300	
$v/m \equiv v^* \rightarrow t$	4.555 <i>T</i>	50	40	39.5	30	20	17.5
250 260 270 280 290	2382.72 2428.27 2473.82 2519.37 2564.92	9.206 8.502 7.489	8.017 7.530	7.958 7.476 7.048	7.134		
300 310 320	2610.47 2656.02 2701.57	6.685 5.987	6.690 6.008	6.654 6.322 6.000	6.723 6.019	6.006	6.003
330 340	2747.12 2792.67	5.423	5.421	5.724	5.414	5.415	5.681 5.402
350 360 380 400 420	2838.22 2883.77 2974.87 3065.97 3157.07	4.926 4.492 4.105 3.777	4.929 4.491 4.112 3.776		4.929 4.489 4.102 3.770	4.913 4.475 4.096 3.755	5.136 4.890 4.387 4.069 3.740,
440 460	3248.17 3339.27	3.468 3.193	3.480 3.200		3.468 3.195	3.459 3.193	3.454 3.193
$egin{array}{l} t_{\mathtt{ont}} & ightarrow \ P_{\mathfrak{s}} & ightarrow \ \delta(v^{*} + \end{array}$	-1)→	250.00 47.654 8.988	263.05 61.054 8.320		280.19 84.008 7.556	304.64 90.477 6.560	312.70 101.05 6.240
$v/m \equiv v^* \rightarrow t$	4.555 <i>T</i>	15	12.5	$ \begin{array}{c} 10 \\ -\delta(v^*+1) \end{array} $	7.5	6.25	
330 340 350 360 370	2747.12 2792.67 2838.22 2883.77 2929.32	5.655 5.367 5.105 4.860	5.303 5.044 4.806 4.586	4.930 4.701 4.489	4.490 4.295	4.126	

Table 37—(Continued)

460

 $t_{\text{sat}} \rightarrow$

 $\delta(v^*+1) \rightarrow$

3339.27

			ubic or (001111111111111111111111111111111111111		
$v/m \equiv v^* \rightarrow$	4.555 <i>T</i>	15	12.5	$ \begin{array}{c} 10 \\ -\delta(v^*+1) \end{array} $	7.5	6.25
380	2974.87	4,427	4.395	4.293	4.114	3.958
400	3065.97	4.049	4.009	3.935	3.784	3.651
420	3157.07	3.723	3.686	3.621	3.491	3.378
440	3248.17	3.432	3.397	3.340	3.226	3.127
460	3339.27	3.173	3.141	3.088	2.988	2.900
$t_{\rm sat} \rightarrow$		321.56	332.03	343.89	357.11	364.05
$P \rightarrow$		113.74	130.32	151.34	177.98	193.49
$\delta(v^* +$	1)→	5.922	5.522	5.076	4.495	4.229
$v/m \equiv v^* \rightarrow$		5.0	4.0	3.0	2.0	
ŧ	4.555 <i>T</i>		δ(v*	+ 1)		_
370	2929.32				2.549	
375	2952.10		3.511	3.103		
380	2974.87	3.723	3.447	3.055	2.460	
390	3020.42	2 440	3.327	2.957	2.370	
400	3065.97	3.449	3.210	2.861	2.280	
410	3111.52		3.100^{a}	2.767		
420	3157.07	3.201	2.990	2.674		
430	3202.62	0.070	2.886			
440	3248.17	2.972	2.785			

From column L of Table 33, $t = 200 \,^{\circ}C$, 4.555T = 2154.97.

2.764 370.53

209.09

3.860

$$P \rightarrow \qquad \qquad 1 \qquad 2 \qquad 3 \qquad 5 \qquad 10 \\ v^* \equiv v/m \rightarrow \qquad 2146 \qquad 1068 \qquad 708.0 \qquad 420.2 \qquad 203.6 \\ \delta(v^* + 1) \rightarrow \qquad 9.0 \qquad 9.4 \qquad 10.2 \qquad 10.2 \qquad 11.3$$

373.40

216.41

3.530

(Cont'd from p. 84)

In the reduction of his observations on the pressures developed by the explosion of mixtures of H_2 and air, D. M. Newitt ¹⁸⁰ used for water-vapor the equation of state (4) proposed by A. Wohl, ^{190a} in which v^* is the specific volume (v/m).

$$P = RT/(v^* - b) - a/Tv^*(v^* - b) + c/T^2v^{*3}$$
 (4)

He gives reasons for believing that b varies with the temperature. Using two different methods of estimation, he obtained the following two sets of values:

T	273	2500	2700	2900	3100	۰K
105b	160	69.5	68.8	68.1	67.4	liters/g-mole-H ₂ O
10⁵ <i>b</i>	160	59.7	58.0	56.0	54.3	liters/g-mole-H ₂ O

All sets of observations of the density and of the specific heat of watervapor indicate that something of the nature of an association sets in as the condition of saturation is approached (see p. 54).

[&]quot;If the observers' interpolated value is used (see Table 35, note) this becomes 3.098.

Table 38.—Defect of Pressure (kg*/cm²) of Dilated Water-vapor: Isometrics

Derived from the data in Table 36 (Nieuwenburg and Blumendal).

 $p = p_i(1 - \delta)$, where $p_i = 4.7064 Tm/v \text{ kg*/cm}^2$, unit of $m/v = \text{kg/cm}^3$, of $4.7064T = 1 \text{ (cm}^3/\text{g)} \cdot (\text{kg*/cm}^2)$, $4.7064T \text{ (cm}^3/\text{g)} \cdot (\text{kg*/cm}^2) = 4.555T \text{ cm}^3 \cdot \text{atm/g}$, T = 273.1 + t. Hence, these values of δ are strictly comparable with those of Table 37.

Unit of v/	$m \equiv v^* = 1$	cm ³ /g, of	£ 4.7064 <i>T</i>	= 1 (cm ³ /	/g)-(kg*/c	m²); δ is	dimensi	onless.	Temp. = t °C
$v/m \equiv v^*$		10	9	8	7	6	5.5	5	4.5
ŧ	4.7064T				δ($v^* + 1) -$			
350	2932.5	4.92							
360	2979.6	4.58	4.53						
370	3026.6	4.32	4.26	4.19	4.13	4.04			
380	3073.7	4.12	4.11	4.05	3.97	3.87	3.82	3.72	3.61
390	3120.8	3.98	3.95	3.90	3.81	3.72	3.66	3.58	3.48
400	3167.8	3.82	3.83	3.74	3.66	3.58	3.52	3.45	3.36
410	3214.9	3.72	3.70	3.67	3.55	3.45	3.40	3.33	3.24
420	3262.0	3.61	3.59	3.55	3.43	3.31	3.28	3.21	3.13
430	3309.1	3.49	3.48	3.41	3.31	3.22	3.18	3.11	3.03
440	3356.1	3.35	3.35	3.29	3.19	3.10	3.06	3.02	2.94
450	3403.2	3.27	3.28	3.19	3.10	3.01	2.96	2.91	2.85
460	3450.2	3.16	3.16	3.07	2.96	2.89	2.86	2.81	2.76
470	3497.3	3.08	3.07	2.98	2.88	2.82	2.79	2.74	2.67
480	3544.4	2.98	2.97	2.90	2.82	2.74	2.70	2.66	2.60
$v/m \equiv v^*$	→	4	3.5	3.0	2.5	2.2	2.0	1.8	1.6
ŧ	4.7064 T					+ 1)			
380	3073.7	3.46	3.28	3 06	2.80	2.61	2.45	2.24	1.91
390	3120.8	3.34	3.18	2.97	2.71	2.52	2.37	2.15	1.80
400	3167.8	3.23	3.07	2.88	2.63	2.45	2.28	2.06	
410	3214.9	3.13	2.98	2.79	2.55	2.37	2.20	1.98	
420	3262.0	3.02	2.89	2.71	2.48	2.29	2.12	1.90	
430	3309.1	2.93	2.80	2.63	2.40	2.21	2.05		
440	3356.1	2.85	2.72	2.55	2.32	2.14	1.98		
450	3403.2	2.76	2.64	2.48	2.25	2.07			
460	3450.2	2.68	2.56	2.40	2.18	1.99			
470	3497.3	2.60	2.50	2.33	2.12				
480	3544.4	2.52	2.42	2.25	2.05				

15. THERMAL ENERGY OF DILATED WATER-VAPOR

In this section are considered the specific heat (c), the enthalpy or heat content, the entropy increase, the decrease in temperature on adiabatic free expansion (Joule-Thomson effect), and certain related quantities, all intimately related to the thermal energy.

Specific Heat of Dilated Water-vapor.

Data on the specific heat of dilated water-vapor are not entirely concordant, especially in the region adjacent to the state of saturation. The situation is complicated by the existence of several more or less contradictory steam tables involving extended extrapolation and much choice and

Table 39.—Formulas and their Coefficients: Specific Heat, Enthalpy, and Entropy of Dilated Water-vapor

(See also Table 43).

Two sets of formulas are given. I. Those by which F. G. Keyes, L. B. Smith, and H. T. Gerry ¹⁹⁷ sum up the results of their program of research on the properties of steam; these supersede all others of the same type that they have published from time to time, and are to be used when the most precise values are desired. II. Those derived by J. Havliček and L. Miškovsky ¹⁹¹ from their own observations. The values defined by the two sets of formulas agree closely, as is shown by the values in Table 40.

 c_p = specific heat at constant pressure, $c_{p\to 0}$ = limit approached by c_p as p approaches zero, p = pressure, H = enthalpy (heat content), S = entropy. Both H and S are measured from saturated water at 0 °C.

I. Keyes, Smith, and Gerry.

Unit of energy = 1 Int. joule, of mass = 1 g; of $t = 1 \text{ kg*/cm}^2$; temp. = t °C Int. scale; $T \equiv 273.16 + t$

1. Limiting value of the specific heat at constant pressure as the pressure is indefinitely reduced $(c_{p\to 0})$.

$$c_{p \to 0} = 1.47198 + 7.5566(10^{-4})T + 47.8365/T$$

2. Value of c_p for pressure p and temperature t. $c_p = c_{p \to 0} + A'p + B'p^2 + C'p^4 + D'p^{13}$, A', B', C', and D' having the following values:

ŧ	10ªA'	а	10bB'	ь	10°C'	с	10dD'	
100	1.5954	1	2.921	2				
120	1.1093	ī	1.446	2	4.596	5		
140	7.979		7.612	3	1.492	5 5		
160	5.907	2 2 2	4.221	2 2 3 3 3	5.306	6		
180	4.483	2	2.447	3	2.035	6		
200	3.477	2	1.474	3	8.320	7		
220	2,748	2	9.183	4	3.595	7	2.812	21
240	2.208	2 2 2 2 2	5.890	4	1.628	7	1.330	22
260	1.801	2	3.877	4	7.683	8	7.416	24
280	1.488	2	2.611	4	3.756	8	4.744	25
300	1,244	2	1.794	4	1.892	8	3.431	26
320	1.051	2	1.255	4	9.787	9	2.765	27
340	8.967	2 2 3 3 3	8.928	4 5 5 5	5.175	9	2.449	28
360	7.71 7	3	6.444	5	2.787	9	2.348	29
380	6.693	3	4.713	5	1.524	9	2.387	30
400	5.848	3	3.489	5	8.423	10	2.485	31
420	5.143	3 3 3 3 3	2.612	55555	4.692	10	2.415	32
440	4.550	3	1.974	5	2.622	10	1.637	33
460	4.049	3	1.506	5	1.463	10	-1.563	34
480	3.622	3	1.159		8.087	11	-1.226	34
500	3.255	3	8.794	6	4.390	11	-4.900	35

3. Value of the enthalpy (H) of dilated water-vapor as measured from saturated water at 0 °C; that is, H is the increase in the "heat content" on changing 1 g of saturated water at 0 °C into dilated water-vapor at temperature t and pressure p.

$$H = 2502.36 + \int_{275.16}^{T} c_{p \to 0} dT - (Ap + Bp^{3} + Cp^{4} + Dp^{18})$$

Table 39—(Continued)

A, B, C, and D having the following values:

t	A	10 <i>bB</i>	ь	10°C	c	10^dD	
100	12.1691	9.1692	1	2.6285	3		
150	6.8776	2.2182	1	1.8411	4		
200	4,4208	6.9814	2	2.0664	5	4.2871	19
250	3.0994	2,6285	2	3.1568	6	2.1742	22
300	2.3103	1.1242	2	5.9107	7	2.7131	25
350	1.8009	5.2816	3	1.2560	7	6.4847	28
400	1.4519	2.6635	3	2.8219	8	2.0549	30
450	1.2013	1.4184	3	6.0829	9	-8.7526	33
500	1.0143	7.8789	4	9.6158	10	-9.9275	34
550	0.8705	4.5225	4	-1.1283	10		

If unit of p=1 atm, these coefficients must be increased numerically by the following amounts: A by 3.323, 3.322, 3.321, and 3.320 per cent, B 6.755 per cent, C 14.00 to 13.97 per cent, and D 52.95 per cent; the first three values for A refer, respectively, to 100, 150, and 200 °C, the fourth to all the others; the first for C refers to 100 °C, the other to all the others. All have been derived from the values of the coefficients as published.

4. Value S of the excess of the entropy of expanded water-vapor at temperature t and pressure p above that of saturated water at 0 °C is given by the formula

$$S = 6.8158 + \int_{-22.10}^{T} (c_{p \to 0}) \frac{dT}{T} - 1.06242 \log_{10} p - (Kp + Lp^2 + Mp^4 + Np^{18})$$

K, L, M, and N having the following values:

91
2
8
3
6

II. J. Havliček and L. Miškovský. 181

Each formula is given in duplicate, first with the Int. steam calorie (= 4.1860 Int. joules), and secondly with the Int. joule as the unit of energy.

Unit of mass = 1 g, of
$$p = 1$$
 kg*/cm²; temp. = t °C (Int. scale); $T \equiv 273.2 + t$.

 $c_{p \to 0} = 0.4402 + 0.0095(t/100) + 0.00072(t/100)^2$ (Int. cal.)

= $1.8427 + 0.03977(t/100) + 0.0030139(t/100)^2$ (Int. joule)

 $H = 597.6 + \int_0^t c_{p \to 0} dt - \{d(p/10^6) + e(p/10^6)^2 + f(p/10^6)^5\}$ (Int. cal.)

= $2501.6 + \int_0^t c_{p \to 0} dt - \{d(p/10^6) + e(p/10^6)^2 + f(p/10^6)^5\}$ (Int. joule)

 $d = 716.64(100/T)^2 + 107.73(100/\tau)^2(3 + 440/\tau) - 1.026$ (Int. cal.)

= $2999.86(100/T)^2 + 450.96(100/\tau)^2(3 + 440/\tau) - 4.295$ (Int. joule)

Table 39—(Continued)

$e = 2.7981(10^7)(100/T)^8 - 0.0726$	(Int. cal.)
$= 11.7128(10^7)(100/T)^8 - 0.3039$	(Int. joule)
$f = 3.1242(10^{18})(100/T)^{22} - 3.8952(10^{17})(100/T)^{21}$	(Int. cal.)
= $13.0779(10^{18})(100/T)^{22} - 16.3053(10^{17})(100/T)^{21}$	(Int. joule)
$\tau \equiv T - 220 = 53.2 + t$	

judgment in smoothing and reconciling the various sets of related data, and also by the repeated publication, with much emphasis on their supposed accuracy, of certain values computed by A. Leduc in 1913, which appear without a literature reference under the designation "best" in his contribution to the *International Critical Tables*.²⁰⁰ These values of Leduc's assume that the general equation of state set up by him for normal gases is applicable to water-vapor, and that the values he used for certain auxiliary data for water-vapor are correct. It may without difficulty be shown that his equation does not satisfactorily represent the best experimental values for water-vapor, and that the auxiliary values he used do not accord with the best data now available. Therefore, whatever his computed data for the specific heat of water-vapor may have been worth in 1913, they can scarcely be accepted now. The more important steam tables will be found listed in Table 259; but little use is made of them in this section.

Data for the specific heat at high temperatures have either been computed from basic constants and spectroscopic data, or based upon the specific heat at constant volume, as inferred from the pressures generated when suitable mixtures of gases are exploded in a closed vessel, and upon the value of $c_p - c_v = T(\delta v/\delta T)_p(\delta p/\delta T)_v$ as computed from an assumed equation of state. Until recently they have been quite uncertain. The interpretation of the spectroscopic data has not been entirely clear. ²⁰¹ In the explosion experiments the temperature is never uniform throughout the volume, the heat losses are difficult to determine, as are the effect of radiation and its absorption, and especially the dissociation; moreover there is not equilibrium between the molecules and the temperature of their immediate surroundings. ²⁰²

In addition to the papers listed elsewhere in this section the following should be examined by those especially interested in this subject.

(1) Determinations by the explosion method:

Langen, A., Mitt. Forsch. Geb. Ing., 8, 1-54 (1903); Bjerrum, N., Z. Elektroch., 17, 731-735 (1911); Siegel, W., Z. physik. Chem., 87, 641-668 (1914); Gallina, V., Ann. d. R. Scuola d'Ing. (Padova), 4, 77-87 (1928); David, W. T., and I.cah, A. S., Phil. Mag. (7), 18, 307-321 (1934); Lewis, B., and von Elbe, G., J. Chem'l Phys., 2, 659-664, 890 (L) (1934); 3, 63-71 (1935); J. Am. Chem. Soc., 57, 612-614 (1935); Phil. Mag. (7), 20, 44-65 (1935); Schmidt, F. A. F., Forsch. Gebiete Ingenieurw., 8, 91-99 (1937).

²⁰⁰ Leduc, A., Int. Crit. Tables, 5, 82 (1929).

²⁰¹ Bonhoeffer, K. F., and Reichardt, H., Z. physik. Chem. (A), 139, 75-97 (1928); Justi, E., Forsch. Gebiete Ingenieurw., 2, 117-124 (1931); Gordon, A. R., and Barnes, C., J. Phys'l Chem., 36, 1143-1151 (1932).

³⁰² Chaudron, G., Bull. Soc. Chim. France (4), 37, 657-679 (1925); McCrea, W. H., Proc. Cambridge Phil. Soc., 23, 942-950 (1927); Bonhoeffer, K. F., and Reichardt, H., loc. cit.

Table 40.—Values of $c_{p\to 0}$ and Its Integrals: Dilated Water-vapor.

By definition, $c_{p\to 0}$ is the limit approached by the specific heat at constant pressure as the pressure is reduced.

I. The following values have been computed by means of the formulas given in Table 39, and are designated as: I = Keyes, Smith, and Gerry's formula $c_{p\to 0}=1.47198+7.5566(10^{-4})T+47.8365/T$, T=273.16+t. II = Havliček and Miškovský's $c_{p\to 0}=1.8427+0.03977(t/100)+0.0030139(t/100)^2$, T=273.2+t.

	Un	it of $c_p = 1$ In	it. joule/g·°C.	Temp. = t °C (Int. scale)	
Form→	1	11	I	II	1	11
<i>t</i>	c _p .	• 0	$\int_{0}^{t}c_{r}$	$\rightarrow 0$ dt —	$\int_0^t (c_z)$	$(t) \rightarrow 0$ $\frac{dt}{T}$
0	1.85352	1.8427	0	0	0	0
50	1.86421	1.8634	92,904	92.64	0.312300	0.31136
100	1.88216	1.8855	186.537	186.36	0.581600	0.58093
150	1.90479	1.9092	281.195	281.21	0.819589	0.81943
200	1.93063	1.9343	377.068	377.29	1.033684	1.03400
250	1.95875	1.9610	474.294	474.68	1.228970	1.23000
300	1.98855	1.9892	572.970	573.42	1.409065	1.40983
350	2.01964	2.0189	673.169	673.61	1.576637	1.57740
400	2.05172	2.0501	774.948	775.32	1.733709	1.73438
450	2.08459	2.0828	878.352	878.64	1.881851	1.88240
500	2.11810	2.1170	983,416	983.62	2.022305	2.02275
550	2.15212	2.1526	1090.168	1090.35	2.156070	2.15650

II. Adapted from tables by E. Justi and H. Lüder²⁰³; conversion to joules and to grams was made by the compiler.

Unit of			indicated.	1 cal = 4.186 j	oules. Temp	$p. = t ^{\circ}C$
Units→ {	g	-mole	g	g-m	ole —	g
f remmo	cal	joule	joule	cal	joule	joule
					C' dt	
t		rp→0			$\int_{0}^{\infty} (c_{p \to 0}) \frac{dt}{T}$	
20	7.98	33.41	1.855	-	-	
100	8.10	33.90	1.882	2.53	10.59	0.588
200	8.32	34.78	1.930	4.48	18.75	1.041
300	8.56	35.83	1.989	6.08	25.45	1.413
400	8.84	37.01	2.054	7.48	31.31	1.738
500	9.12	38.18	2.120	8.68	36.33	2.017
600	9.41	39.39	2.186	9.81	41.06	2.279
70 0	9.72	40.69	2.259	10.85	45.42	2.521
800	10.02	41.94	2.328	11.82	49.48	2.746
1000	10.58	44.28	2.458	13.58	56.84	3.155
1200	11.08	46.38	2.574	15.14	63.38	3.518
1400	11.52	48.22	2.678	16.58	69.40	3.852
1600	11.88	49.72	2.761	17.90	74.93	4.159
1800	12.19	51.03	2.832	19.13	80.08	4.445
2000	12.45	52.12	2.894	20.28	84.89	4.712
2500	12.95	54.21	3.009	22.80	95.44	5.298
3000	13.23	55.38	3.074	24.98	104.57	5.804

²⁰³ Justi, E., and Lüder, H., Forsch. Gebiete Ingenieurw., 6, 209-216 (1935). Superseding Justi, E., Idem, 5, 130-137 (1934).

(2) Computation from fundamental constants and spectroscopic data:

Henning, F., and Justi, E., Wiss. Abh. d. Phys. Tech. Reichs., 14, 171-174 (1930-31) = Z. techn. Physik, 11, 191-194 (1930); Hausen, H., Forsch. Gebiete Ingenieurw., 2, 319-326 (1931); Zeise, H., Z. Elektroch., 39, 758-773, 895-909 (1933); Trautz, M., and Ader, H., Z. Physik, 89, 12-14 (1934); Lewis, B., and von Elbe, G., J. Chem'l Phys., 3, 63-71 (1935); J. Am. Chem. Soc., 37, 612-614 (1935); Wilson, E. B., Jr., J. Chem'l Phys., 4, 526-528 (1936); Kassel, L. S., Chem. Rev., 18, 277-313 (1936); Murphy, G. M., J. Chem'l Phys., 5, 637-641 (1937).

(3) Reviews, summaries, etc.:

Callendar, H. L., Phil. Trans. (A), 215, 383-399 (1915); World Power, 1, 274-280, 325-328 (1924); 3, 302-312 (1925); Fischer, V. Z. techn. Physik, 5, 17-21, 39-44, 83-88 (1924); Jazyna, W., Idem, 6, 261-262 (1925); Z. Physik, 57, 341-344 (1929); Plank, R., Z. techn. Physik, 5, 397-404 (1924); Saunders, S. W., J. Phys'l Chem. 28, 1151-1166 (1924); Davis, H. N., and Keenan, J. H., Proc. World Eng. Cong. (Tokyo), 4, 239-264 (1931).

(4) Steam tables, see Section 90.

Table 41.—Specific Heat of Dilated Water-vapor at Constant Pressure: Preferred Values

Various other sets of values will be found in Tables 42 and 43.

The following values are those given by Keyes, Smith, and Gerry ¹⁹⁷ and computed by means of the formula $c_p = c_{p\to 0} + A'p + B'p^2 + C'p^4 + D'p^{13}$ in which $c_{p\to 0}$ and the several coefficients take the values given in Tables 39 and 40, respectively; values for other values of t and P may be similarly computed.

	Unit of	$c_p = 1$ Int. jou	le/g = 0.23907	calis/g; of P =	= 1 kg*/cm2.	Temp. = t °C.	
$t \rightarrow$	340	360	380	400	420	440	460
<i>P</i> 120	5.708	4.479	3.836	3.427	3,152	2,960	2.822
140	8.948	5.624	4.501	3.877	3.474	3.200	3.007
160 180		7.791	5.420 6.864	4.441 5.168	3.862 4.332	3.481 3.810	3.218 3.459
200				6.164	4.905	4.195	3.735

With the preceding values Keyes, Smith, and Gerry ¹⁹⁷ compare those published by W. Koch, ²⁰⁴ finding the following differences: $\Delta = \text{Koch} - \text{KSG}$; unit of $\Delta = 1$ Int. joule/g.

$\stackrel{t o}{P}$	340	360	380	400 1000Δ	420	440	460
120 140 160 180 200	+ 80 + 21	- 13 + 34 - 2	- 23 - 52 - 34 - 46	+ 5 - 43 - 68 - 58 - 37	+ 37 - 5 - 41 - 59 - 47	+ 62 + 31 + 1 - 22 - 27	+ 78 + 56 + 38 + 19 + 6

Table 42.—Specific Heat of Dilated Water-vapor at Constant Pressure: Various Sets of Values

(See also Tables 41, 43, and 55)

Preferred values are given in Table 41; those here given indicate values in use in 1932. They serve to show how the values run, and are useful in making rough approximations; many of them differ by several units in the second decimal place from the preferred values defined by the formula used in Table 41 (see bottom of that table).

Table 42—(Continued)

1 joule/g = 0.9997 Int. joule/g = 0.2390 cal₁₈/g = 18.015 joule/g·mole = 4.305 cal₁₈/g-mole. 1 kg/cm² = 0.96784 atm = 0.98066 bar.

Unit of P=1 kg*/cm², of $P_a=1$ atm, of $P_b=1$ bar = 10° dynes/cm², of $c_p=1$ joule/g·°C, of $C_p=1$ cal/g-mole·°C. Temp. = t°C.

T.	Ω	Knoblauch	A bne	Winkhaus.206

$P_a P_b t_{ m sat}$		0.5 0.4839 0.4903 0.9	1.0 0.9678 0.9807 99.1	1	2.0 1.9357 1.9613 19.6		6.0 5.807 5.884 58.1		10 9.678 9.807 79.1
		2 000	0.020*		c _p - 2.097*		2.323		2.565
t_{ext} 110		2.000 1.971	2.0384		2.097		2.323		2.303
120		1.963	2.021 2.009						
140		1.950	1.988		2.055				
160							2.310		
		1.946	1.971		2.026				
180		1.946	1.971		2.005		2.222		
200		1.954	1.971		1.996		2.151		2.381
220		1.958	1.971		1.996		2.105		2.260
240		1.967	1.975		1.996		2.072		2.185
260		1.975	1.980		1.996		2.059		2.143
280		1.984	1.988		2.005		2.055		2.122
300		1.992	2.000	•	2.013		2.055		2.109
320		2,000	2.009		2.021		2.059		2.105
340		2,013	2.017		2.030		2.063		2.105
360		2.026	2.030		2.038		2.072		2.109
380		2.034	2.038		2.051		2.080		2.118
P	1:	2	14		16		18		20
P_a		1.614	13.550		15.485		17,421		19.357
P_b		1.768	13.729		15.691		17.652		19,613
tent		7.1	194.2	2	00.5		06.2		11.4
t					c				
toat		2.687	2.808		2.925		3.051		3.180
200		2.528	2.712						
220		2.360	2.482		2,620		2.787		2.984
240		2.264	2.348		2.440		2.544		2.678
260		2,201	2.260		2.331		2,402		2,490
280		2.164	2.214		2.264		2.318		2.377
300		2.147	2.184		2.226		2.268		2.310
320		2.147 2.138	2.168		2.201		2.235		2.268
			2.100						
340		2.134	2.159		2.184		2.214		2.239
360		2.130	2.151		2.176		2.197		2.222
380		2.134	2.151		2.168		2.189		2.210
II. O.	Knoblau	ich and W	7. Koch.206						
P	19.	9	29.9			40.1		60	.2
P_a P_b	19. 19.	260 515	28.93 29.33			38.810 39.325		58	.264 .036
- •	t	Ср	t	C _P			Cp	1	Cp.
	332.9	2.277	241.6	3.353	252	.8 .3	.881	280.3	4.625
	334.8	2.260	246.6	3.219	257		.805	283.8	4.353

²⁰⁴ Koch, W., Forsch. Gebiete Ingenieurw., 3, 1-10 (1932).

²⁰⁵ Knoblauch, O., and Winkhaus, A., Mitt. Forsch. Geb. Ing., 195, 1-20 (1917) \rightarrow Z. Ver. deuts. Ing., 59, 376-379, 400-405 (1915).

²⁰⁰ Knoblauch, O., and Koch, W., Z. Ver. Deut. Ing., 72, 1733-1739 (1928) \rightarrow Mech. Eng., 51, 147-150 (1929).

Table 42—(Continued)

$P \rightarrow$	1	9.9	29.9		40.1		60.2	
		Cp	t	Cp	1	Cp	t	Cp
	335.9	2.244	259.3 283.8 320.1 353.9 358.4 386.3	2.968 2.671 2.449 2.378 2.382 2.315	263.8 274.0 288.5 322.7 371.2 403.7	3.495 3.261 2.989 2.650 2.491 2.411	287.5 301.1 324.6 334.6 368.0 369.2	4.161 3.679 3.156 3.051 2.742 2.683
			433.9 437.0 493.2	2.327 2.340 2.344	436.5 436.8 494.0	2.390 2.390 2.382	429.0 437.7	2.532 2.537
$P_a P_b$		80.0 77.427 78.453			00.2 96.978 98.263		120.1 116.2 117.2	238
	297.6 300.5 309.7 326.2 341.5 375.0 377.5 421.0	5.6 5.3 4.7 3.8 3.4 3.0 3.0	7572 603 701 881 666 918 930 733	314.9 317.9 318.0 324.5 335.0 352.5 376.1 411.9 410.1	6.39 6.38 5.95 5.66 4.72 3.95 3.42 3.03 3.06	8 32 2 33 8 34 2 34 1 35 8 36 0 38	9.6 8.6 5.4 2.2 5.1 2.7 5.9 9.0 7.6	8.234 7.828 6.488 5.706 5.542 5.023 4.295 3.608 3.110
III. W	. Koch.	04						
$P_{oldsymbol{a}} \ P_{oldsymbol{b}} \ t_{oldsymbol{sat}}$		120 116.14 117.68 323.11	130 125.8 127.4 329.2	9	140 135.50 137.29 335.04	150 145.13 147.10 340.5)	160 154.85 156.91 345.71
t t _{sat} 340 360		8.625 5.791 4.468	9.74 7.00 4.9	09	11.020 8.973 5.661	12.50		14.307 7.792
380 400 420		3.814 3.433 3.190	4.1 3.6 3.3	22	4.451 3.835 3.471	4.8 4.0 3.6	86	5.389 4.375 3.823
440 460		3.023 2.902	3.1: 2.9	24	3.232 3.065	3.3. 3.1.	54	3.484 3.257
$P \\ P_a \\ P_b \\ t_{\mathtt{eat}}$		170 164.53 166.71 350.66		180 174.21 176.52 355.38	Ср	190 183.89 186.33 359.88	1	00 93.57 96.13 64.18
t_{mat} 360		16.560 9.730 6.025		19.428 13.511 6.821		23.121 22.828 7.863		27.885 9.383
380 400 420		4.714 4.036		5.112 4.275		5.577 4.551		6.130 4.861
440 460		3.630 3.362		3.789 3.479		3.969 3.605		4.170 3.743

²⁰⁷ Trautz, M., and Steyer, H., Forsch. Gebiete Ingenieurw., 2, 45-52 (1931).

²⁰⁶ Jakob, M., Z. Ver. Deuts. Ing., 56, 1980-1988 (1912). ²⁰⁶ Knoblauch, O., and Mollier, H., Mitt. Forsch. Geb. Ing., 108 and 109, 79-106 (1911) → Z. Ver. Deuts. Ing., 55, 665-673 (1911).

Table 42—(Continued)

IV. M. Trautz and H. Steyer.207

$P_{oldsymbol{a}\ P_{oldsymbol{b}}}$	150 145.176 147.100	200 193.568 196.133	250 241.960 245.166	300 290.352 294.200
t			·	
350	8.79			
400	4.18	7.32	9.67	14.35
450	3.10	4.14	4.39	5.19
500	2.51	3.35	4.18	4.98

V. From compilation by A. Leduc ²⁰⁰ based on observations by M. Jakob ²⁰⁸ and by O. Knoblauch and H. Mollier. ²⁰⁹

$P_a P_b$	1.033 1 1.013	2.066 2 2.026	4.133 4 4.053	6.199 6 6.080	10.333 10 10.132	14.465 14 14.186	20.665 20 20.265
<i>t</i> 100	2.017			р			
120	1.996	2.093					
140	1.984	2.046					
160	1.975	2.021	2.155	2.356			
180	1.971	2.005	2.101	2,226	2.645		
200	1.971	1.996	2.067	2.151	2.381	2.808	
250	1.980	1.996	2.034	2.072	2.155	2.264	2.461
300	1.996	2.009	2.034	2.059	2.109	2,159	2.247
350	2.021	2.025	2.051	2.067	2.109	2.147	2.214
400	2.051	2.059	2.076	2.093	2.122	2.155	2.205
450	2.084	2.093	2.113	2.118	2.139	2.164	2.201
500	2.122	2.126	2.134	2.143	2.159	2.176	2.201
550	2.156	2.159	2.164	2.168	2.180	2.189	2.210

VI. Derived from c_v , as inferred from the pressures developed when confined mixtures of H_2 and O_2 are exploded, by means of an assumed equation of state: $c_p - c_v = T(\delta v/\delta T)_p^2(\delta p/\delta T)_v$. Pressure = 1 atm.

Ref. b→	ICT	HH	E	Ref.b→	ICT	нн	E
t		c _p		1		ср	
100	2.040		1.966	1000	2.297	2.335	2.462
200	2.010	1.946	1.996	1200	2.485	2.544	2.636
300	2.004		2.032	1400	2.708	2.796	2.837
400	2.010	1.980	2.074	1600	2.977		3.067
500	2.030		2.122	1800	3.289		3.320
600	2.062	2.055	2.178	2000	3.470		3.601
700	2.103		2.238	2200	3.586		3.902
800	2.158	2.172	2.305	2300	3.614		4.064
900	2.221		2.382				2.002

VII. Computed by Gordon and Barnes 210 from basic constants, Steinwehr's equation of state, 211 and spectroscopic data. Pressure = 1 atm.

\boldsymbol{T}	ŧ	C_p	Cp	\boldsymbol{T}	t	C_{p}	C _P
400	126.9	8.26	1.920	900	626.9	9.46	2.199
500	226.9	8.45	1.964	1000	726.9	9.74	2.264
600	326.9	8.68	2.017	1100	826.9	10.02	2.329
700	426,9	8.93	2.076	1200	926.9	10.29	2.392
800	526.9	9.19	2.136		7-017	10.10	2.072

 ²⁰⁰ Gordon, A. R., and Barnes, C., J. Phys'l Chem., 36, 1143-1151 (1932).
 ²¹¹ Steinwehr, H. v., Z. Physik, 3, 466-476 (1920).

Table 42—(Continucd)

VIII. A. Leduc.²¹² Computed by him from certain other data for water-vapor, assuming the validity of the general equation of state set up by him as applicable to all normal gases. In his compilation,³⁰⁰ these values are called the "best," and are given without citation. They have been republished.²¹⁸ It will be noticed that they disagree with the experimental values in magnitude and in that the latter pass through a minimum as the temperature is decreased, the pressure remaining constant; whereas Leduc's computed values continually decrease.⁶

$t \rightarrow P_a$	100	120	140	150	160
1	1.81,	1.833	1.87,	1.93	2.00,
2 3		1.871	1.91 ₈ 1.95 ₉	1.95,	2.02
4				2.026	2.07
$P_{\mathtt{ext}}$	1.81	1.87,	1.971		

^a For the limiting value of c_p at t_{sat} , the following values by M. Jakob ²⁰⁸ are given in the compilation by A. Leduc. ²⁰⁰

^b References:

E = E. D. Eastman.²¹⁴

HH = L. Holborn and F. Henning.215

ICT = adapted from the compilation by A. Leduc, 200 apparently derived directly from Table C (p. 205) of Partington and Shilling's "The Specific Heat of Gases" (London, 1924). There, two sets of values are given. In each of them the same value of c_{ν} is used, but in one set $c_{\nu} - c_{\nu}$ is said to have been derived from Berthelot's equation of state; and in the other, from Callendar's. The former is the smaller by 5.8, 1.6, 0.5, and 0.05 per cent at 100°, 200°, 300°, and 400 °C, respectively. The average of the two sets is given in ICT, but here those ascribed to Callendar's equation are used. The value of c_{ν} is derived from c_{ν} and $c_{\nu} - c_{\nu}$.

^e Leduc ²¹⁶ maintains that this experimentally observed minimum arises from experimental errors, and that his computed values are to be preferred. But it can be shown that the subsidiary data employed by him were not sufficiently accurate, that the test he employed to demonstrate the applicability of his equation of state to water-vapor was not sufficient, and that that equation does not satisfactorily represent the specific volume of water-vapor. Whatever may have been the worth of these computed values in 1913, one is scarcely justified in regarding them more highly than the corresponding experimental data available today. Additional information regarding his equation and his method of computation may be found in the Comptes Rendus.²⁰⁷

Table 43.—Specific Heats of Water-vapor at 1 atm: Comparison of Interpolation Formulas

(See also Table 39)

Callendar's formula is $c_p = 1.997_9 + 38.49(373.1/T)^{10/3}P_{atm}/T$ joules/g.°C; KSG's is that given in Table 39; the others are of the form $c = A + B(t/100) + C(t/100)^2 + D(t/100)^3$, the coefficients having the values

²¹² Leduc, A., Ann. de chim. et phys. (8), 28, 577-613 (1913).

²¹⁸ Leduc, A., Jour. de Phys. (6), 2, 24-30 (1921).

²¹⁴ Eastman, E. D., U. S. Bureau of Mines Tech. Paper 445, (1929).

Table 43—(Continued)

given at the heads of the several columns. Eastman's formula (E) is not supposed to give the details of the actual variations, but merely values that are always near the true value. Each tabulated value of the specific heat has been computed by means of the indicated formula.

		Te	emp. = t °C; un	it of $c=1$ jou	ile/g·°C		
Ref.a→	KSG	Ca	R	HSH	Sh	E	F
A			1.96	1.8948	2.2292	1.942	2.311
B C			- 0.015	+ 0.0386		+ 0.0205	- 0.745
D			+ 0.0051	0	$+0.0343_{\circ}$ -0.00118_{\circ}	+ 0.00311	+ 0.46 ₀ 0
1				cp	- 0.001100		(cp) satb
100	2.078	2.101	1.952	1.933	2.0475	1.96₅	2.026
120		2.08_{0}	1.951	1.9412	2.018 ₈	1.971	2.07,
140		2.064	1.951	1.9490	1.9928	1.97,	2.16,
160		2.052	1.951	1.9567	1.968	1.983	2.29,
180		2.042	1.95 _t	1.964	1.946,	1.98,	2.462
200	1.968	2.03,	1.952	1.9722	1.927	1.998	2.661
250		2.022	1.95₅	1.991	1.888	2.013	3.322
300	2.002	2.014	1.961	2.010_{8}	1.862	2.032	4.216
350		2.00,	1.970	2.0302	1.8477	2.51,	5.45°
400	2.058	2.006	1.982	2.049	1.8442	2.074	
450		2.004	1.996	2.068,	1.850,	2.097	
500	2.122	2.00.	2.018	2.0882	1.8667	2.122	
550		2.001	2.032	2.1078	1.891	2.14,	
600		2.000	2.053	2.1268	1.922	2.177	
700		1.99	2.105	2.165	2,0048	2.238	
800		1.99	2.16	2.2042	2.1061	2.305	
900		1.99	2.238	2.242,	2.2197	2.380	
1000		1.99	2.320	2.2816	2.338,	2.46	
1200		1.998	2.515	2.3590	2.562,	2.64	
1400		1.99	2.751	2.4363	2.7212	2.84	
1600		1.998	3.030	2.5137	2.7598	3.07	
1800		1.99	3.34,	2.5910	2.618	3.32	
2000		1.998	3.70,	2.6684	2.243	3.60	
2200		1.998	4.110	2.7458	1.5767	3.90	
2300		1.998	4.327	2.7844	1.116	4.06	
Ref."→	KSG	Ca	R	HSH	Sh	E	Sh
A			1,957	1.914	2.132	1.95	1.687
\boldsymbol{B}			-0.00623	+0.01934		+0.0112	-0.177_{2}
C D			$+0.0017_{2}$	0	$+0.0111_{6}$	+ 0.001037	
υ			0	0	- 0.00029 ₅	0	- 0.000931
t			Mean c	between 100			Co
100	2.078	2.10 ₁	1.95	1.93,	2.048	1.96₺	1.539
120 140		2.09 ₀ 2.08 ₀	1.952	1.937	2.03 ₃ 2.01 ₉	1.968	1.516.
140		2.000	1.951	1.941	2.019	1.97	1.495

²¹⁸ Holborn, L., and Henning, F., Ann. d. Physik (4), 23, 809-845 (1907).

²¹⁶Leduc, A., Chem. Rev., 6, 1-16 (1929).

²¹⁷ Leduc, A., Compt. rend., 152, 1752-1756 (1911); 153, 51-54 (1911); 154, 812-815 (1912).

			Table 43	–(Continue	(d)		
Refa→	KSG	Ca	R	HSH	Sh	E	Sh
t			Mean cp betv	veen 100 and	t °C		Co
160		2.073	1.95,	1.94	2.00_{6}	1.974	1.476
180		2.067	1.95	1.94	1.994	1.977	1.458
200	1.994	2.06	1.952	1.95	1.98_{2}	1.98₀	1.443
250	1.987	2.05_{0}	1.953	1.962	1.95_{7}	1.988	1.414
300	1.988	2.042	1.953	1.972	1.93₅	1.996	1.395
350	1.993	2.036	1.956	1.982	1.920	2.00₅	1.387
400	2.002	2.03_{1}	1.95,	1.99_{1}	1.90_{7}	2.015	1.388
450	2.012	2.027	1.964	2.00_{1}	1.89,	2.02₅	1.399
500	2.023	2.024	1.96,	2.01_{1}	1.894	2.036	1.418
550	2.036	2.022	1.975	2.02_{0}	1.89_{2}	2.04_{7}	1.444
600		2.02_{0}	1.982	2.03_{0}	1.893	2.05_{8}	1.478
700		2.016	1.99_{7}	2.04,	1.904	2.08_a	1.563
800		2.014	2.017	2.069	1.926	2.11_{0}	1.668
900		2.012	2.04_{0}	2.08_{8}	1.955	2.13,	1.787
1000		2.01 ₀	2.067	2.107	1.99_{1}	2.17	1.914
1200		2.00_{8}	2.13_{0}	2.14	2.07₅	2.24	2.170
1400		2.00_{7}	2.20_{7}	2.185	2.16_{3}	2.31	2.393
1600		2.00_{6}	2.29_{7}	2.22_{3}	2.24,	2.40	2.538
1800		2.00_{s}	2.40_{2}	2.26_{2}	2.29_{5}	2.49	2.559
2000		2.004	2.52_{0}	2.30_{1}	2.311	2.59	2.412
2200		2.00_{3}	2.652	2.339	2.275	2.70	2.053
2300		2.00_{3}	2.724	2.35,	2.232	2.76	1.779

a References:

Ca = H. L. Callendar. 218

= E. D. Eastman.214

F = V. Fischer.210

HSH = L. Holborn, K. Scheel, and F. Henning. 200

= M. Randall.221 Sh = W. G. Shilling. 222

Table 44.—Mean Internal Specific Heat of Dilated Water-vapor at Constant Pressure 223

By definition $\bar{c}_{ip} \equiv \bar{c}_p - p\Delta v/\Delta t$, where \bar{c}_p is the mean specific heat at constant pressure over the range t to $(t + \Delta t)$, Δv is the accompanying increase in the specific volume, and p is the pressure expressed in appropriate units. Were the vapor ideal, \bar{c}_{ip} would be \bar{c}_{v} , the specific heat at constant volume, and would be independent of p.

b At the pressure of the saturated vapor, i.e., the value that would be found at that pressure by extrapolating the curve defined by observations taken at slightly lower pressures and at the temperature indicated.

At critical temperature (374 °C) the equation gives $(c_p)_{sat} = 5.95_s$ joules/g·°C.

gas Callendar, H. L., "Properties of Steam," pp. 98, 60, 61, 15 (1920).

²¹⁹ Fischer, V., Z. Physik, 43, 131-151 (1927).
220 Holborn, L., Scheel, K., and Henning, F., "Wärmetabellen" (1919).
221 Randall, M., Int. Crit. Tables, 7, 231 (1930).

²²² Shilling, W. G., Phil. Mag. (7), 3, 273-301 (1927).

Derived from Callendar, H. L., Proc. Inst. Mech. Eng., 1929, 507-527 (1929).

Table 44—(Continued)

Unit of	p = 1 (1b*/in	2)La; of F	$P_a = 1$ atm	= 1.01325	bars; of cip	= 1 joule/	g.°C. Ten	np. = t °C
	$t \rightarrow$		$t_{\mathtt{sat}}$	250	300	350	400	450
Þ	Pa	trat			- číp between			
400	27.232	228.8	1.819	1.787	1.782	1.695	1.669	1.644
450	30.636	235.2	1.847	1.821	1.753	1.711	1.677	1.652
500	34.040	241.2	1.880	1.864	1.781	1.733	1.697	1.674
600	40.848	251.9	1.941	1.952	1.840	1.772	1.730	1.694
700	47.656	261.2	2.000		1.901	1.819	1.766	1.724
800	54.464	269.7	2.064		1.967	1.863	1.797	1.650
900	61.272	277.4	2.137		2.044	1.914	1.834	1.780
1000	68.080	284.5	2.200		2.132	1.967	1.877	1.816
1200	81.696	297.2	2.347		2.346	2.087	1.958	1.880
1400	95.312	308.4	2.511			2.231	2.051	1.941
1600	108.928	318.4	2.691			2.405	2.158	2.030
1800	122.544	327.5	2.889			2.622	2.278	2.306
2000	136.160	335.8	3.114			2.899	2.417	2.204
2400	163.392	350.6	3.665				2.776	2.424
2800	190,624	361.3	4.394				3.273	2.706
3200	217.856	373.6	5.476				4.055	3.084
3600	245.088	3.0					5.334	3.584
4000	272.320						0.000	4.314

"For g = 981.16 cm/sec2, value at London.

Table 45.—Specific Heat of Dilated Water-vapor at Constant Volume

I. Derived from data in Table 47 where remarks and references are given. Pressure is very low.

	nit of $c_v =$	1 joule/	g.°C; of	P = 1 kg* emp. == t°	$C = T \circ K$	$P_a = 1$ at	m; of Pb	= 1 bar.	
Source	ea→ T	PS	$Ch\mathbf{K}$	КЈ	M	c, J	Misc.	T_A	Тв
0 12.3 100.0 104.56	273.1 285.4 373.1 377.66	1.530	1.391 1.419	1.410	1,421	1.415	Br ^a 1.570	1.368	1.394
104.36 107.5 140.0 160.0 180.0 190.0	380.6 413.1 433.1 453.1 463.1			1.412 1.417 1.422 1.426			1.570	1.403	1.433
200.0 202.6 210.0 220.0 230.0 240.0	473.1 475.7 483.1 493.1 503.1 513.1	1.532	1.473	1.426 1.431 1.438 1.442 1.452	1.471	1.452		1.442	1.475
250.0 260.0 270.0 280.0 290.0	523.1 533.1 543.1 553.1 563.1			1.456 1.463 1.473 1.480 1.517					4 700
297.8 300.0 310.0 320.0 330.0 340.0 350.0	570.9 573.1 583.1 593.1 603.1 613.1 623.1	1.536	1.533	1.505 1.494 1.531 1.542 1.561 1.575		1.514		1.484	1.528

For reference see Table 47.

			T	able 45	(Continue	d)			
S	$ \begin{array}{c} \text{ource}^a \rightarrow \\ T \end{array} $	PS	ChK		M	J	Misc.	TA	Тв
360.0	633.1			1.594		C#			
370.0	643.1			1.610					
380.0	653.1			1.626					
390.0	663.1			1.647					
392.9	666.0							1.528	1.594
400.0	673.1	1.545		1.668	1.591	1.606	\mathbf{K}^{a}		
410.0	683.1						1.621		
488.1	761.2							1.577	1.645
500.0	773.1	1.567					1.656		
526.9	800.0		1.680						
550.0	823.1						1.689	4 404	4 740
583.2	856.3	4 500			4 504		He	1.624	1.710
600.0	873.1	1.599			1.721		1.668		
650.0	923.1						1.686	1 650	1 777
678.4	951.5 973.1	1.640					1.705	1.659	1.777
700.0 750.0	1023.1	1.040					1.726		
773.9	1047.0						Wa	1.684	1.842
800.0	1073.1	1.696			1.853		2.030	1.001	1.012
850.0	1123.1	1.070			1.000		2.716		
868.9	1142						2.7.10	1.749	1.905
900.0	1173.1	1.759							
926.9	1200		1.944						
963.9	1237							1.824	1.963
1000.0	1273.1	1.836			1.978				
1058.9	1332							1.902	2.014
1153.9	1427							1.977	2.065
1200.0	1473.1	2.023			2.090				
1248.9	1522						P^a	2.046	2.128
1250.0	1523.1		0.162				2.416		
1326.9	1600		2.163				Ba		
1340.0	1613.1 1617						2.335	2.132	2.193
1343.9 1400.0	1673.1	2.247			2.185			2.102	2.170
1439.9	1713	2.221			2.103			2.230	2.267
1534.9	1808							2.337	2.353
1540.0	1813.1						2.530	2.001	21000
1600.0	1873.1	2.516							
1629.9	1903							2.432	2.432
1726.9	2000		2.323						
1800.0	2073.1	2.809							
1819.9	2093							2.644	
2000.0	2273.1	3.008							
2200.0	2473.1	3.127							
2300.0	2573.1	3.152						0 760	
Lit	nit							2.7 69	
					of Table 4				$v = T \times$
		, as give	-	nde's equ	ation of s	-			
$P \rightarrow$	0.5	1	2	4	$P \rightarrow$	0.5	1	2 .	4
$P_a \rightarrow$	0.484	0.968	1.936	3.871	P_a		0.968	1.936	3.871
$P_b \rightarrow$	0.490	0.981	1.961	3.923	$P_b \rightarrow$	0.490	0.981	1.961	3.923
*	4.405	C.			1,00	4 470	4 404	1 506	4 574
110	1.487	1.513			160	1.470	1.481	1.506	1.571
120	1.481 1.477	1.506 1.497	1 522		170	1.472	1.484	1.499 1.495	1.557
130 140	1.471	1.497	1.532 1.523		180 190	1.473 1.477	1.486 1.488	1.493	1.542 1.530
150	1.469	1.487	1.512	1.588	200	1.482	1.490	1.494	1.520
	faranca car	Table 47	012	2.000	200	1.302	1.270	1.777	4.020

2.396

2.457

Table 46.—Mean Specific Heat of Dilated Water-vapor at Constant Volume

(See also Table 51)

Derived from columns SS and M of Table 48, which see. The fourth digit of \bar{c}_v is uncertain by several units.

Uni	it of $\bar{c}v = 1$ jo	oule/g. °C = 0.	2390 cal ₁₅ /g.	C. Temp.	$= t \circ C = T$	°K; range 0 t	to t°C
R	ef.a→	SS	M	Re	f. a→	SS	M
t	T	č		ı	T	Č1	
0	273.1	1.434	1.470	1600	1873	1.790	1.775
100	373	1.452	1.480	1700	1973	1.826	1.814
200	473	1.471	1.486	1800	2073	1.865	1.856
300	573	1.492	1.496	1900	2173	1.907	1.905
400	673	1.510	1.504	2000	2273	1.951	1.956
500	773	1.529	1.516	2100	2373	1.998	2.008
600	873	1.547	1.529	2200	2473	2.044	2.060
700	973	1.565	1.543	2300	2573	2.092	2.113
800	1073	1.586	1.564	2400	2673	2.141	2.170
900	1173	1.605	1.589	2500	2773	2.192	2.222
1000	1273	1.624	1.608	2600	2873	2.246	2.277
1100	1373	1.645	1.631	2700	2973	2.301	2.338

2800

2900

3000

3073

3173

3273

1673 1773 "For references see Table 48.

1473

1573

1200

1300

1400

1500

Table 47.—Molecular Specific Heat of Dilated Water-vapor at Constant Volume 224

1,656

1,686

1.715

1.742

1.670

1,698

1.726

1.759

(See also Table 45)

The several values are said to have been derived from the indicated sources. Those from PS are the same as those given as "best" by A. Leduc ²⁰⁰ and are defined by the formulas $C_v = 6.750 - 0.00119t +$ $(2.34(t/1000)^2)$ if $(100 \, ^{\circ}\text{C}) = (1700 \, ^{\circ}\text{C})$, and $(1000)^2$ if 100 $4.67(t/1000)^2$ if $1700 \,^{\circ}\text{C} \gtrsim t \gtrsim 2300 \,^{\circ}\text{C}$. Trautz gives two sets of smoothed values: T_A , based on all the observations; T_B , based on only those from spectroscopic data. He thinks the B values are to be preferred to about 1200 °C. He states that the observed values have as far as possible been reduced to infinite volume.

Unit of $C_v = 1 \text{ cal/gfw-H}^2\text{O} \cdot \text{°C} = 0.05551 \text{ cal/g} \cdot \text{°C} = 0.2323 \text{ joule/g} \cdot \text{°C}$. Temp. = $t \cdot \text{°C} = T \cdot \text{K}$. Pressure very low

Sourc	e ^a → T	PS	ChK	KJ	M	J	Misc.	TA	Тв
0 12.3	273.1 285.4		5.99			,		5.89	6.00
100.0 104.56	373.1 377.66	$6.58_{\scriptscriptstyle 5}$	6.11	6.07	6.11 ₈	6.09	Br ^a 6.76	0.07	0.00
107.5	380.6						0.70	6.04	6.17
140.0	413.1			6.08					
160.0	433.1			6.10					
180.0	453.1			6.12					

²²⁴ Trautz, M., Ann. d. Physik (5), 9, 465-485 (1931).

Table 47.—(Continued)

				77. (00	, in conca,				
	rce ^a →	PS	ChK	KJ	M	J	Misc.	T_{Λ}	Тв
t 190.0 200.0 202.6	T 463.1 473.1 475.7	6.598	6.34	6.14 6.14	6.33 ₁	6.25		6.21	6.35
210.0 220.0	483.1 493.1			6.16 6.19				0.21	0.55
230.0 240.0 250.0	503.1 513.1 523.1			6.21 6.25 6.27					
260.0 270.0 280.0	533.1 543.1 553.1			6.30 6.34 6.37					
290.0 297.8 300.0	563.1 570.9 573.1	6.612	6.60	6.53 6.48		6.52		6.39	6.58
310.0 320.0 330.0 340.0 350.0	583.1 593.1 603.1 613.1 623.1			6.43 6.59 6.64 6.72 6.78					
360.0 370.0 380.0 390.0	633.1 643.1 653.1 663.1			6.86 6.93 7.00 7.09					
392.9 400.0 410.0	666.0 673.1 683.1	6.652		7.18	6.84,	6.91	Ka 6.98	6.58	6.86
488.1 500.0 526.9	761.2 773.1 800.0	6.74.	7.23				7.13	6.79	7.08
550.0 583.2 600.0	823.1 856.3 873.1	6.88₅	,,,,,		7.408		7.27 Ha 7.18	6.99	7.36
650.0 678.4 700.0	923.1 951.5 973.1	7.062			7.108		7.26 7.34	7.14	7.65
750.0 773.9 800.0	1023.1 1047 1073.1	7.30			7.97 ₆		7.43 Wa 8.74	7.25	7.93
850.0 868.9	1123.1 1142				1.916		11.69	7.53	8.20
900.0 926.9 963.9	1173.1 1200 1237	7.57:	8.37		0.74			7.85	8.45
1000.0 1058.9 1153.9	1273.1 1332 1427	7.902			8.51₅			8.19 8.51	8.67 8.89
1200.0 1248.9 1250.0 1326.9	1473.1 1522 1523.1 1600	8.708	9.31		8.998		Pa 10.4 Ba	8 81	9.16
1340.0 1343.9 1400.0	1613.1 1617 1673.1	9.67:			9.40 _a		10.05	9.18	9.44
1439.9 1534.9 1540.0	1713 1808 1813.1	J.013).10g		10.89	9.60 10.06	9.76 10.13
1070.0	1010.1						10.03		

			Table 4	7.—(Cor	itinued)				
Sou t	$T^{\text{rce}^a \to}$	PS	ChK	KJ	M C_v -	J	Misc.	TA	Тв
1600.0 1629.9 1726.9	1873.1 1903 2000	10.83	10.0					10.47	10.47
1800.0 1819.9	2073.1 2093	12.096						11.38	
2000.0 2200.0	2273.1 2473.1	12.95 13.46							
2300.0 Lir	2573.1 nit	13.57						11.92	

" Sources:

- = Bjerrum, N., Z. physik. Chem., 79, 513-536 (1912).
- =Brinkworth, J. H., Phil. Trans. (A), 215, 383-438 (1915).
- ChK = Chemikerkalender, 1931; derived from spectroscopic data.
- = Holborn, L., and Henning, F., Ann. d. Physik (4), 18, 739-756 (1905); 23, 809-845 (1907).
- = Jakob, Max, private communication to Trautz.
- Jakob, Max, private communication to Trautz.

 Knoblauch and associates: Knoblauch, O., and Jakob, M., Sitz. K. Bayer (Münch.)

 Akad. Wiss. (Math.-phys.), 35, 441-446 (1905); Mitt Forsch. Geb. Ing., 35,

 36, 109-152 (1906) → Z. Ver. Deuts. Ing., 51, 81, 88, 124-131 (1907); Knoblauch, O.,

 and Mollier, H., Mitt. Forsch. Geb. Ing., 108, 109, 79-106 (1911) → Z. Ver. Deuts.

 Ing., 55, 665-673 (1911); Knoblauch, O., and Raisch, E., Idem, 66, 418-423 (1922);

 Knoblauch, O., and Winkhaus, A., Mitt. Forsch. Geb. Ing., 195, 1-20 (1917) → Z. Ver.

 Deuts. Ing., 59, 376-379, 400-405 (1915). K
- KJ = Knoblauch, O., and Jakob, M., Mitt. Forsch. Geb. Ing., 35, 36, 109-152 (1906) → Z. Ver. Deuts. Ing., 51, 81-88, 124-131 (1907).
- = Mecke, R., unpublished communication to Trautz; from band spectrum.
- Misc = Several sources: B, Br, H, K, P, and W.
- =Pier, M., Z. Elektroch., 15, 536-540 (1909); 16, 897-903 (1910).
- PS =Partington, J. R., and Shilling, W. G., "The Specific Heat of Gases," London, Benn, 1924.
- =Trautz, M., Ann. d. Physik (5), 9, 465-485 (1931); see head of table for significance of A and B.
- =Womersley, W. D., Proc. Roy. Soc. (London) (A), 100, 483-498 (1921); 103, 183-184 (1923). Criticism by R. T. Glazebrook, Idem, 101, 112-114 (1922).
- ^b The published value (11.84) seems to have been computed by means of the formula valid below 1700 °C. This and several other values belonging in the PS column were assigned by Trautz to ChK.

Table 48.—Mean Molecular Specific Heat of Dilated Water-vapor at Constant Volume

The mean molecular specific heat at constant volume over the indicated range in temperature is \tilde{C}_v . The values at high temperatures have been inferred from the pressures developed on exploding mixtures of H_2 and O_2 , initial pressure being 1 atm; those at lower temperatures, from C_p and an assumed equation of state. Comments on the several series will be found with the appropriate references.

Unit of $C_v = 1$ cal/gfw-H₂O.°C = 0.05551 cal/g.°C = 0.2323 joule/g.°C. Temp. = t°C = T°K

Range→			0 to t °C						
	f.°→ T	CG	NW	SS	M	w	100 to t °C		
΄0	273.1	6.00	5.99	6.17	6.33				
100	373.1	6.05	6.05	6.25	6.37	6.59	6.65		
200	473.1		6.14	6.33	6.40	6.61	6.63		
300	573.1		6.24	6.42	6.44	6.65	6.61		
400	673.1			6.50	6.47	6.70	6.62		

Table 48.—(Continued)

_	_			. (00///////	,		
t F	$Range \rightarrow Ref.^a \rightarrow T$	CG	NW	— 0 to t °C —	- CvM	w	100 to t °C PS
500 527 600	773.1 800.1 873.1	6.32	6.53	6.58 6.66	6.53 6.58	6.76 6.85	6.63 6.64 6.67
700 800	973.1 1073.1			6.74 6.83	6.64 6.73	6.92 7.02	6.72 6.78
900 927	1173.1 1200.1		7.09	6.91	6.84	7.14	6.86 6.89
1000 1100 1200	1273.1 1373.1 1473.1	6.82		6.99 7.08 7.19	6.92 7.02 7.13	7.29 7.50 7.75	6.96 7.07 7.20
1300 1327	1573.1 1600.1		7.61	7.31	7.26	8.03	7.34 7.40
1400 1500 1600	1673.1 1773.1 1873.1	7. 50	,,,,,	7.43 7.57 7.71	7.38 7.50 7.64	8.35 8.74 9.23	7.50 7.68 7.87
1700 1727 1800	1973.1 2000.1 2073.1		8.10	7.86 8.03	7.81 7.99	9.82 10.46	8.07 8.13 8.29
1900 2000	2173.1 2273.1	8.36		8.21 8.40	8.20 8.42	11.07 11.69	8.51 8.74
2100 2127	2373.1 2400.1	8.55	8.50	8.60	8.64		8.96 8.98
2200 2300 2400	2473.1 2573.1 2673.1	8.75 8.96 9.17		8.80 9.01 9.22	8.87 9.10 9.34		9.17 9.36
2500 2527	2773.1 2800.1	9.39	8.8	9.44	9.57		
2600 2700 2800	2873.1 2973.1 3073.1	9.62 9.85 10.09		9.67 9.91	9.80 10.06 10.31		
2900 3000 3100 3200 3300	3173.1 3273.1 3373.1 3473.1 3573.1	10.34 10.60 10.86 11.13 11.41			10.58 10.85		
3400 3500 3600 3700 3800	3673.1 3773.1 3873.1 3973.1 4073.1	11.70 11.99 12.29 12.59 12.91					
3900 4000	4173.1 4273.1	13.23 13.56	_				

Ran	ıge→	18 to t °C	Ran	ıge→	15 to	1°C	16 to t °C
Re	f."→	WM	Re	$f_{\cdot}a \rightarrow$	В	M W	N
1	T	Cv	ı	T		C,	
1758	2031	8.26	1400	1673		10.7	
1770	2043	8.20	1750	2023		10.6	
1781	2054	8.07^{b}	1811	2084	7.92d		
1882	2155	8.32	1950	2223		10.8	
1973	2246	8.38^{b}	2110	2383	8.54d	10,0	
2035	2308	8.54^{b}	2120	2393		10.6	
2060	2333	8.546	2377	2650	9.37d	10.0	
2092	2365	8.70	2663	2936	10.0		

Table 48 (Continu	ued)
-------------------	------

ge→ ^a → T	18 to t°C WM		$a \rightarrow$	B 15 t	MW — Cu	16 to t °C N
2377 2421	8.72 8.65	2908 3060	3181 3333	10.5 10.9		
2455 2584 2591 2603 2677	8.58° 8.78 8.73 8.76 8.66°	2327 2427 2527 2627 2727	2600 2700 2800 2900 3000			10.25 10.32 10.40 10.44 10.50
	T 2377 2421 2455 2584 2591 2603	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

SS ^a	Range→		273.1 to t°		
t	T	Cv	1	T	Co
- 273.1	0	5.96	1327	1600	7.11
- 173	100	5.96	1427	1700	7.23
- 73	200	5.98	1527	1800	7.35
+ 27	300	6.03	1627	1900	7.49
127	400	6.09	1727	2000	7.64
227	500	6.16	1827	2100	7.80
327	600	6.23	1927	2200	7.98
427	700	6.31	2027	2300	8.16
527	800	6.39	2127	2400	8.35
627	900	6.47	2227	2500	8.55
727	1000	6.56	2327	2600	8.75
827	1100	6.64	2427	2700	8.96
927	1200	6.73	2527	2800	9.17
1027	1300	6.82	2627	2900	9.40
1127	1400	6.92	2727	3000	9.61
1227	1500	7.01			

" References and notes:

- = N. Bjerrum.²²⁵ Those marked ^d are from observations by Pier. These values are quoted
- by B. Neumann ²²⁰ and from him by CG.

 A. D. Crow and W. E. Grimshaw, ²²⁷ values are defined by the formula: $C_v = 6 + (7/15) \cdot (t/1000) + (16/45) \cdot (t/1000)^2$, between 0 °C and t °C.
- = Muraour,228 values are derived from the observations by Pier and Bjerrum. of mean Cr given in a compilation by A. Leduc 200 were taken from this list.

- NW = W. Nernst and K. Wohl.281
- PS = Computed by the compiler from the formulas for C_ν given by Partington and Shilling, see Table 47.
- SS =F. Schmidt and H. Schnell.232
- = W. D. Womersley.^{23t} Values for t < 1000 were derived from earlier data obtained by others.
- WM = K. Wohl and M. Magat.284
- b, c See 284.
- " See " B.

²²⁵ Bjerrum, N., Z. Elektroch., 18, 101-104 (1912).

²²⁶ Neumann, B., Z. angew. Chem., 32, 141-146 (1919).

²²⁷ Crow, A. D., and Grimshaw, W. E., Phil. Trans. (A), 230, 39-73 (1931).

²²⁸ Muraour, Chim. ct indus., 10, 23-29 (1923).

²²⁹ Maxwell, G. B., and Wheeler, R. V., J. Chem. Soc., 1928, 15-21 (1928).

²⁸⁰ Newitt, D. M., Proc. Roy. Soc. (London) (A), 125, 119-134 (1929).

²³¹ Nernst, W., and Wohl, K., Z. techn. Phys., 10, 608-614 (1929).

²⁸² Schmidt, F., and Schnell, H., Z. techn. Phys., 9, 81-92 (1928); computed from available data. ²⁸³ Womersley, W. D., *Proc. Roy. Soc. (London) (A)*, 100, 483-498 (1921); 103, 183-184 (1923); see criticism by Glazebrook, R. T., *Idem*, 101, 112-114 (1922).

Table 49.—Ratio of the Principal Specific Heats of Dilated Water-vapor

 $\gamma = c_p/c$

Unit of $P=1 \text{ kg*/cm}^2$; of $P_a=1 \text{ atm}$; of $P_b=1 \text{ bar}$. Temp. $=t \, ^{\circ}\text{C} = T \, ^{\circ}\text{K}$

I. From PS of Table 45 and ICT of Section VI of Table 42. Pressure = 1 atm. Except for the change noted in Table 47, note b, these are the ones contained in the table of "best" values for the molecular specific heat in A. Leduc's compilation ²⁰⁰ and derived from Partington and Shilling, "The Specific Heat of Gases."

	γ	t	γ	1	γ
100	1.334	700	1.283	1600	1.183
200	1.311	800	1.272	1800	1.171
300	1.305	900	1.263	2000	1.154
400	1.301	1000	1.251	2200	1.147
500	1.295	1200	1.228	2300	1.146
600	1.290	1400	1.205		

II. From the velocity of sound. W. G. Shilling.²²² Equation of state used was either Callendar's (Cal) or Berthelot's (Ber).

Eq.→	Cal.	Ber.	Eq.→	Cal.	Ber.
£		y 	t		γ
100	1.332	1.317	600	1.316	1.316
200	1.334	1.332	700	1.295	1.296
300	1.337	1.337	800	1.277	1.277
400	1.335	1.336	900	1.257	1.257
500	1.329	1.330	1000	1.241	1.241

III. From Section II of Table 45 and section I of Table 42.

\boldsymbol{P}		0.5	1	2	4
P_a		0.484	0.968	1.936	3.871
$P_{m a} \\ P_{m b}$		0.490	0.981	1.961	3.923
	T			γ	
110	• 383	1.325	1.336		
120	393	1.325	1.336		
130	403	1.325	1.333	1.352	
140	413	1.325	1.332	1.349	
150	423	1.325	1.331	1.348	1.376
160	433	1.324	1.331	1.345	1.372
170	443	1.322	1.328	1.343	1.368
180	453	1.321	1.326	1.341	1.365
190	463	1.320	1.325	1.339	1.362
200	473	1.318	1.323	1.336	1.360

IV. Values computed by A. Leduc ²³⁵ by his method, employing his equation of state for normal gases; and in his compilation ²⁰⁰ marked "best." See remarks in Section VIII of Table 42.

$P_a^{t o}$	120	130	140	150	160
1	1.365		1.364	1.333	1.314
2		1.37	1.36	1.344	1.326
3			1.37	1.356	1.34
4				1.37	1.35
$P_{\mathtt{sat}}$	1.3784		1.380		

²²⁴ Wohl, K., and Magat, M., Z. physik. Chem. (B), 19, 117-138 (1932); derived from 3 sets (unmarked, b, and c) by Wohl, K., and von Elbe, G., Idem, 5, 241-271 (1929).

Table 49 .- (Continued)

V. Derived by H. G. Muhammad ²⁸⁶ from his determinations of γ for air saturated with water-vapor at the temperature indicated. Total pressure about 745 mm-Hg.

t	γ	t	γ	t	γ
15	1.3421	35	1.3723	55	1.3054
20	1.3568	40	1.3671	60	1.3015
25	1.3672	45	1.3406	65	1.2993
30	1.3736	50	1.3199	70	1.2976
For 100 °	C and Past he gives	v = 1.373.			

Table 50.—Ratio of the Principal Specific Heats of Air Saturated with Water-vapor ²³⁶

P = tc	otal pressure, $\gamma = c_i$	ρ/c_v for the saturated air.	Unit of P	= 1 mm-Hg.	Temp. = t °C.
t	P	γ	t	P	γ
15	742.1	1.4009	45	745.6	1.3949
20	743.4	1.4008	50	745.4	1.3893
25	744.3	1.4007	55	745.0	1.3828
30	741.8	1.4005	60	745.0	1.3765
35	743.3	1.4000	65	745.0	1.3697
40	745.1	1.3989	70	745.0	1.3618

Table 51.—Enthalpy of Dilated Water-vapor

Several sets of the better values are compared in Table 52. The enthalpy ("heat-content," "total heat") of dilated water-vapor as measured from saturated water at 0 °C is H; $c_p = (\delta H/\delta t)_p$; $\bar{c}_p = (H_t - H_{t_1})/(t - t_1)$ is the mean value of c_p over the indicated range; P_s kg*/cm² is the saturation pressure at the indicated temperature.

An earlier and more detailed table, in which the unit of pressure is 1 lb*/in², has been published by H. L. Callendar, ¹⁹² and very detailed tables and graphs may be found in the numerous steam tables (see Table 259).

Unit of $P_a = 1$ atm, of P = 1 kg*/cm², of H = 1 Int. joule/g.°C. Temp. = t°C (Int. scale)

I. Keyes, Smith, and Gerry.

Based on their formula given in Table 39; computed and published by themselves. Very detailed tables on the same basis, but in terms of °F and $lb*/in^2$, have been published by J. H. Keenan and F. G. Keyes, "Thermodynamic Properties of Steam," 1936. Only the last few digits of the value of H for P are given here, the others being the same as for the corresponding value of P_a . For example, the value of H for P=25 and t=300 is 3010.53.

Leduc, A., Ann. d. chim. ct phys. (8), 28, 577-613 (1913); J. de Phys. (6), 2, 24-30 (1921).
 Muhammad, H. G., Bull. Acad. Sci., (Allahabad), 3, 269-294 (1934).

T 11		· · ·	7 1
Table	51	(Continue	ea)

			Table Ji.	Comme	4)		
	<i>t</i> →	100	150	200	250	300	200 300
P_{\bullet}	P			H			Ĉp
1		2675.35	2776.22	2874.79	2973.43	3072.94	1.9815
	1	5.80	6.46	4.94	3.54	3.02	1.9808
5	_			2854.71	2959.95	3063.11	2.0840
•	5			5.58	60.50	3.53	2.0795
10	-			2826.07	2941.79	3050.26	2.2419
10	10			8.06	4.01	1.11	2.2305
25	10			0.00	2877.66	3008.14	2,2000
25	25				81.51	10.53	
50	25				01.01	2921.77	
30	EO					8.04	
	50	0675.25	0744.04	0700 47	0700.00	2747.05	
P_{sat}	P_{sat}	2675.35	2744.84	2790.46	2799.00	2/4/.05	
	$t\rightarrow$	350	400	450	500	550	450
Pa	P			н			5 5 0 Ĉp
1		3173.67	3275.81	3379.48	3484.72	3591.58	2.1210
•	1	3.73	5.86	9.52	4.76	1.61	2.1210
5	•	3166.10	3269.74	3374.48	3480.51	3587.97	2.1349
~	5	6.40	70.14	4.68	0.68	8.12	211017
10	•	3156.37	3262.04	3368.16	3475.21	3583,44	2.1528
	10	7.00	2.52	8.57	5.55	3.74	
25		3125.45	3238.02	3348.74	3459.05	3569.70	2.2096
	25	7.17	9.34	9.80	9.91	70.43	
50		3067.51	3194.99	3314.84	3431.26	3546.30	2.3146
	50	71.51	7.87	7.08	3.03	7.83	
100		2918.67	3095.64	3240.77	3372.45	3497.70	2.5693
	100	30.01	102.67	5.80	6.36	500.80	
150		2677.85	2971.96	3156.96	3309.40	3446.65	2.8969
	150	710.39	85.28	65.64	15.21	51.67	
200			2809.41	3060.81	3240.77	3393.09	3.3228
	200		33.55	74.19	9.86	400.11	
250			2550.33	2948.90	3166.89	3336.95	3.8805
-	250		607.05ª	68.32	79.21	46.15	
P_{sat}	$P_{\it eat}$	2562.58					

II. Havliček and Miškovský. (See also Table 52.)

Based on their formula given in Table 39. Conversion into joules was made by the compiler.

100	150	200	250	00	20
		Н			30 č
2674.0	2775.3	2874.1	2972.9 2882.5	3072.5 3008.5	1.9
350	400	450 //	500	2927.3 550	45 55
3173.4	3275.5	3378.9	3484.0	3591.2	2.1
3125.3 3068.8	3237.9 3195.6	3348.8 3315.7	3459.3 3432.1	3569.8 3547.6	2.2 2.3
2928.1	3099.3	3242.9	3374.3	3500.3	2.5

Table 51	(Continued)
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$t \rightarrow$	350	400	450	500	550	450 550
P			H			CP
150	2707.5	2981.7	3162.1	3312.4	3450.9	3.888
200		2828.5	3070.4	3245.8	3398.6	3.282
250		2608.7	2964.5	3173.8	3343.8	3.793
300			2838.1	3096.8	3286.8	4.487
350				3013.1		
400				2922.2		

III. W. Koch.²³⁷ Conversion into joules was made by the compiler.

<i>t</i> →	200	225	250	275	300	325	200
P				H			300 Čp
1	2871.6	2920.6	2970.4	3020.6	3070.4	3121.1	1.988
5	2852.8	2903.8	2956.2	3008.9	3060.4	3111.4	2.076
10	2826.0	2882.5	2938.2	2993.0	3046.6	3099.3	2.206
25		2804.6	2878.3	2943.6	3004.3	3063.7	
50				2836.4	2922.7	2998.0	
75					2809.2	2920.1	
100						2818.8	
$\stackrel{i\rightarrow}{P}$	350	375	400	425 —— H	450	475	500
1	3171.7	3222.8	3273.9	3325.8	3378.1	3430.4	3483.6
5	3163.8	3216.1	3266.8	3320.3	3372.7	3426.6	3479.4
10	3153.3	3206.9	3258.8	3313.2	3366.8	3421.2	3474.8
25	3121.9	3179.7	3234.5	3292.7	3348.8	3405.3	3461.0
50	3066.2	3130.3	3191.4	3254.2	3315.3	3375.2	3434.2
75	3004.3	3078.8	3147.4	3214.8	3281.0	3345.0	3407.4
100	2930.2	3020.6	3100.2	3174.2	3245.0	3313.2	3378.9
125	2834.8	2952.0	3045.3	3129.4	3207.3	3278.5	3349.2
150	2701.2	2870.3	2983.8	3079.6	3164.2	3242.5	3316.6
175		2768.6	2913.4	3022.7	3116.5	3203.5	3284.3
200		2625.4	2830.2	2961.2	3065.4	3162.5	3250.0
225		2327.4	2729.3	2890.8	3011.4	3119.0	3214.8
250		1856.1	2602.8	2813.0	2956.2	3073.4	3176.8
275		1817.1	2429.1	2726.8	2896.7	3026.0	3137.8
300		1792.9	2194.3	2633.8	2833.9	2976.2	3096.0

IV. Third International Steam-Table Conference, 1934. For allowed tolerances, etc., see Table 260.

$t\rightarrow$	100	150	200	250	300	350	400	450	500	550
-	1.03323	4.8535	15.857	40.560	87.611	168.63				
P						H				
1	2676	2777	2875	2973	3074	3174	3276	3380	3485	3592
5			2855	2959	3063	3166	3269	3374	3481	3588
10			2827	2940	3048	3155	3270	3369	3476	3584
25				2880	3006	3125	3238	3350	3461	3570
50					2924	3069	3195	3315	3433	3548
75					2816	3005	3149	3280	3405	3525

²²⁷ Koch, W., Z. Ver. deuts. Ing., 78, 1160 (1934).

Table 51.—(Continued)										
t→ Paat→ P	100	150 4.8535	200 15.857	250 40.560	300 87.611	350 168.63	400	450	500	550
100 125 150 200						2929 2834 2709	3099 3044 2982 2833	3243 3204 3163 3071	3375 3346 3316 3249	3501 3477 3452 3400
250 300 Psat	2676	2746	2792	2801	2747	2562	2607 2196	2962 2837	3177 3097	3345

eValues in Section I at 400 °C for P and $P_a = 250$ are not so good as the others, density > 0.1 g/cm².

Table 52.—Enthalpy of Dilated Water-vapor: Comparison of Several Sets of Values

Adapted from a table published by J. Havliček and L. Miškovský. 191 Not checked by the compiler.

The values of II (enthalpy, counted from saturated water at $0 \,^{\circ}$ C) are given for set A; for the other sets only the excess (ΔH) of each value over the corresponding one for A is given. For example, at P = 1 and t = 150 the value of H for A is 663.3; for P it is 663.3 - 0.2 = 663.1.

Some of the values here given for the P set differ slightly from the corresponding ones in Table 51, which Havliček and Miškovský computed by means of their equation.

Unit of $P=1$ kg*/cm², of H and $\Delta H=1$ Int. steam cal/g=4.1860 Int. joule/g. T	$\Gamma \text{emp.} = \iota \circ C \ (I$	nt. scale)
--	---	------------

$\stackrel{t\rightarrow}{P}$	Ref.	150	200	250	300	350 H and ΔH	400	450	500	550
1	A P G I	663.3 - 0.2 - 0.1	686.9 - 0.3 - 0.9 - 0.4	710.4 - 0.1 - 0.8 - 0.3	734.2 - 0.1 - 0.7 - 0.2	758.2 0.0 - 0.5 - 0.2	782.6 0.0 - 0.5 - 0.2	807.3 0.0 - 0.3 - 0.1	832.4 - 0.1 - 0.2 - 0.1	858.0 - 0.4 - 0.2
25	A P G C	- 0.1	- 0.1	688.4 - 0.2 - 0.8 - 1.0 - 0.6	719.2 - 0.9 - 1.5 - 1.5 - 1.2	747.1 - 0.7 - 1.3	773.8 - 0.4 - 1.1 - 0.6 - 0.5	800.2b - 0.2 - 0.2 - 0.4 - 0.2	826.5 - 0.2 + 0.3 0.0 0.0	852.9 - 0.3
50	A P G C				699.5 - 0.5 - 1.3 - 1.4 - 1.1	733.8 - 1.1 - 1.3 - 1.2 - 0.9	763.9 - 0.8 - 1.5 - 1.0 - 0.8	792.4 - 0.5 - 0.2 - 1.1 - 0.8	820.1 -0.3 $+0.3$ -0.4 -0.2	847.4 - 0.1
100	A P G C					700.0 - 0.6 0.0 - 0.9 - 0.5	741.2 - 1.0 - 0.6 - 1.6 - 1.2	775.4 - 0.9 - 0.2 - 1.8 - 0.9	806.5 - 0.5 + 0.7 - 0.8 - 0.5	836.3 - 0.2 - 0.2
150	A P G C I						713.2 - 0.7 - 0.4 - 1.5	756.2 - 1.0 - 0.3 - 1.4	792.0 - 0.7 + 0.3 - 0.2	824.6 - 0.2
	1						- 1.1	- 0.9	- 0.2	- 0.2

Table	52	(Continued)
Table	34.	Commune

$\stackrel{t\rightarrow}{P}$		150	200	250	300	350	400	450	500	550
P	Ref.ª					$-H$ and ΔI	4			
200	Α						676.9	734.3	776.3	812.3
	A P						-0.8	-0.8	-0.9	-0.3
	G						-0.8	-2.0	+ 0.1	
	I						-0.4	- 0.9	-0.3	- 0.3
250	Α						623.3	708.7	759.3	799.5
	P G						-0.7	-0.4	-1.0	- 0.6
	G						-1.5	-2.5	-0.4	
	Ī						- 0.8	-1.2	-0.5	- 0.6
300	A						525.1			
	\mathbf{P}						+ 0.4			
	G						-0.9			
	Ī						- 0.6			

- ^a References: The authors do not give the references more specifically than here.
 - A A.S.M.E.: Davis, Keenan, Keyes, Osborne, Smith. (These seem to be the equivalent of the values in Section I of Table 51.)
 - C B.E.M.A.A.: Egerton, Callendar.
 - G Germany: Hausen, Henning, Jakob, Koch. (This is identical with the set of values published by Koch and given in Table 51.)
 - I Third International Steam Table Conference, 1934. See Section IV of Table 51.
 - P M.A.P.: Havliček, Miškovský.

Table 53.—Isopiestic Variation in the Enthalpy of Water Substance Through the Critical Temperature 191

The following values of J. Havliček and L. Miškovský have been taken from their Table 1. The enthalpy (II) is measured from saturated water at 0 °C.

Since the saturation temperature for P=200 is 364.1 °C, and for P=225 is 373.9 °C, and the critical pressure and temperature are 225.5 kg*/cm² and 374.11 °C, respectively, it is obvious that all values lying above the heavy line refer to the liquid; and all below it, to the vapor.

Unit of P=1 kg*/cm²; of H=1 Int. steam cal/g=4.1860 joule/g. Temp. = t °C (Int. scale)

$P \rightarrow t$	200	225	250	275 H	300	350	400
20	24.55			-	26.79		28.92
100	103.8				105.54		107.22
200	205.5	•			206.0		206.9
300	318.2				316.7		316.2
350	392.6	389.45	387.8	385.4	383.9	381.45	379.3
360	415.6	410.0	406.1	402.9	400.5		
370	610.5	439.5	428.7	422.7	418.4		
375						421.3	416.4
380	640.4	596.5	470.0	448.0	440.7		
390	661.4	630.2	585.7	500.7	470.3		
400	646.1	652.6	622.6	581.6	525.5	478.5	463.8
410					582.2		

^b The value for A at P = 25, t = 450 is printed 802.0, which is evidently too great; the corresponding value in Table 51 leads to 800.2.

Table 55.—(Commun)											
$l^{\prime} \rightarrow l$	200	225	250	275 —— II ——	300	350	400				
450	733.5	720.2	708.3	693.1	677.9	644.5	607.3				
500	775.4	767.5	758.3	749.4	639.8a	719.6	698.6				
550	812.0	804.3	798.9								

[&]quot; So printed; probably should be 739.8.

Table 54.—Entropy of Dilated Water-vapor (See also Table 55)

The excess of the entropy of the vapor at the indicated temperature and pressure above that of saturated water at $0 \,^{\circ}$ C is S; S_s is the value of S at the indicated pressure and the corresponding saturation temperature t_s ; ΔS is the change in S when the temperature is changed over the indicated range while the pressure remains constant at the indicated value.

Unit of
$$P=1$$
 kg*/cm² = 0.96784 atm; of $P_a=1$ atm; of $p=1$ (lb*/in²)L^a; of S , S_a , and $\Delta S=1$ Int. joule/g °C. Temp. = t °C (Int. scale)

I. Preferred values: F. G. Keyes, L. B. Smith, and H. T. Gerry. 197 Based on their formulas in Table 39; conversion from calories to joules (1 Int. steam calorie = 4.1860 Int. joules) was made by the compiler. Very detailed tables on the same basis but in terms of °F and lb*/in² have been published by J. H. Keenan and F. G. Keyes, "Thermodynamic Properties of Steam," 1936.

$t \rightarrow$	100	200	300	400	500	400 500
P			s			Δ.S
1	7.3686	7.8425	8.2221	8.5482	8.8375	0.2893
10		6.7022	7.1313	7.4699	7.7663	0.2964
50			6.2216	6.6570	6.9831	0.3261
100				6.2283	6.6093	0.3810
200				5.5820	6.1630	0.5810
250					5.9889	

II. Adapted from R. Mollier.²³⁸ Not to be preferred to values in I.

$t\rightarrow$		t.	200	300	400	500	200 300	400 500
\boldsymbol{P}	t.			s			,	s —
1	99.1	7.373	7.860	8.241	8.560	8.833	0.382	0.273
2	119.6	7.145	7.531	7.918	8.238	8.512	0.386	0.274
2	132.9	7.012	7.336	7.727	8.049	8.323	0.391	0.275
4	142.9	6.917	7.195	7.591	7.914	8.190	0.395	0.276
6	158.1	6.781	6.992	7.397	7.724	8.001	0.405	0.277
8	169.6	6.684	6.841	7.257	7.587	7.866	0.416	0.279
10	179.0	6.607	6.719	7.147	7.481	7.761	0.426	0.280
20	211.4	6.353		6.790	7.144	7.432		0.288
30	232.8	6.189		6.562	6.939	7.235		0.296
40	249.2	6.061		6.382	6.788	7.093		0.305
50	262.7	5.952		6.222	6.666	6.980		0.314
100	309.5	5.546			6.232	6.610		0.378
150	340.5	5.240			5.882	6.367		0.484
200	364.2	4.905			5.512	6.168		0.656

Table 54.—(Continued)

III. Adapted from H. L. Callendar. 192 Not to be preferred to values in I.

	$t\rightarrow$		t _a	300	400	500	400 500
Þ	P_a	t.					ΔS
400	27.232	228.8	6.252	6.611	6.986	7.296	0.310
500	34.040	241.2	6.168	6.480	6.870	7.176	0.306
600	40.848	251.9	6.096	6.366	6.773	7.085	0.312
800	54.464	269.7	5.974	6.168	6.614	6.938	0.324
1000	68.080	284.5	5.870	5.985	6.482	6.821	0.339
1200	81.696	297.2	5.777	5.802	6.368	6.722	0.354
1600	108.928	318.4	5.604		6.171	6.560	0.389
2000	136.160	335.8	5.437		5.989	6.426	0.437
2400	163.392	350.6	5.259		5.806	6.308	0.502
2800	190.624	363.1	5.057		5.606	6.200	0.594
3200	217.856	373.6	4.789		5.362	6.098	0.736
3600	245.088				5.031	6.000	0.969
4000	272.320					5.901	

[&]quot;Value at London, where g = 981.16 cm/sec².

Table 55.—Various Thermal Data for Dilated Water-vapor: Computed from Spectroscopic Data

G = Gibbs function (H - TS), often called the "free energy at constant pressure," H = enthalpy or heat content, S = entropy, T = absolute temperature, $E_0 = \text{internal energy at 0 °K}$, $C_p = \text{specific heat at constant pressure}$. G, H, and S are measured from 0 °K, *i.e.*, each is the increase in the corresponding property when the temperature is increased from 0 °K to that indicated.

I. A. R. Gordon.²³⁹ It is assumed that the pressure is so low that the vibration of each H_2O molecule is essentially uninfluenced by the presence of neighboring molecules. Giauque has independently computed S for T=298.1, 463.1, and 485.0 °K, using thermal data and obtaining values that are smaller than Gordon's by about 0.1 per cent.²⁴⁰ In order that Gordon's value for R shall be the same as that adopted by the *International Critical Tables* and used elsewhere in this compilation it is necessary to use the relation 1 cal = 4.1873 joules in converting his values from calories to joules.

Temp. = $T \circ K = t \circ C$. 1 cal /g-mole = 0.23243 joule /gram

Unit→		$G-E_0$	cal per g-mole	deg	$G-E_0$ 1 je	oule per gram	deg
T	t	$-\frac{\sigma}{T}$	S	C_{p}	$-\frac{U}{T}$	S	C_{p}
298.1	25.0	37.179	45.101	8.000	8.642	10.483	1.859
300	26.9	37.230	45.151	8.002	8.653	10.494	1.860
350	76.9	38.452	46.389	8.066	8.937	10.782	1.875

²⁰⁸ Molliet, R., "Neue Tabellen und Diagramme," 5th ed., 1927.

²²⁰ Gordon, A. R., J. Chem'l Phys., 2, 65-72, 549(L) (1934); supersedes Idem, 1, 308-312 (1933). ²⁴⁰ Giauque, W. F., and Stout, J. W., J. Am. Chem. Soc., 58, 1144-1150 (1936); Giauque, W. F., and Archibald, R. C., Idem, 59, 561-569 (1937).

Table 55.—(Continued)

Unit \rightarrow 1 cal per g-mole deg		deg	$G-E_0$ 1 joule per gram deg				
T	t	$-\frac{\sigma}{T}$	S	C_p		S	C_p
400	126.9	39.513	47.472	8.155	9.184	11.034	1.895
450	176.9	40.452	48.439	8.260	9.402	11.259	1.920
500	226.9	41.296	49.315	8.379	9.598	11.462	1.948
550	276.9	42.062	50.119	8.504	9.776	11.649	1.976
600	326.9	42.765	50.864	8.635	9.940	11.822	2.007
650	376.9	43.415	51.561	8.771	10.091	11.984	2.039
700	426.9	44.020	52.216	8.910	10.232	12.136	2.071
750	476.9	44.587	52.836	9.053	10.363	12.281	2.104
800	526.9	45.121	53.425	9.199	10.487	12.418	2.138
850	576.9	45.627	53.987	9.347	10.605	12.548	2.172
900	626.9	46.106	54.525	9.497	10.716	12.673	2.207
950	676.9	46.563	55.043	9.648	10.823	12.794	2.242
1000	726.9	46,999	55.542	9.799	10.924	12.910	2.278
1050	776.9	47,418	56.023	9.948	11.021	13.021	2.312
1100	826.9	47,820	56.489	10.095	11.115	13.130	2.346
1150	876.9	48,206	56.941	10.240	11.204	13.235	2.380
1200	926.9	48,579	57.380	10.382	11.291	13.337	2.413
1250	976.9	48.940	57.807	10.522	11.375	13.436	2.446
1300	1026.9	49.289	58.223	10.656	11.456	13.533	2.477
1400	1126.9	49.956	59.022	10.914	11.611	13.718	2.537
1500	1226.9	50.586	59.783	11.153	11.758	13.895	2.592
1750	1476.9	52.03	61.54	11.67	12.093	14.304	2.712
2000	1726.9	53.32	63.13	12.09	12.393	14.673	2.810
2250	1976.9	54.49	64.58	12.4	12.665	15.010	2.88
2500	2226.9	55.57	65.90	12.7	12.916	15.317	2.95
2750	2476.9	56.56	67.12	12.9	13.146	15.601	3.00
3000	2726.9	57.49	68.25	13.1	13.362	15.863	3.04

II. A. R. Gordon and C. Barnes.²⁴¹ The spectroscopic data upon which these values are based have been superseded by those of R. Mecke,²⁴² W. Baumann and R. Mecke,²⁴³ and K. Freudenberg and R. Mecke.²⁴⁴ This fact must be considered by the user. Irrespective of their accuracy, they are of interest in showing the distribution of S among the three types of motion. Subscripts: t = translational, r = rational, v = vibrational entropy. The value of S_t has been corrected for the departure of water-vapor from ideality. Steinwehr's equation of state ²⁴⁵ being used.

Unit of Sa	, S_r , S_v , and S	= 1 cal/g-mole.º	C = 0.23243 joul	e, g.°C. Temp.	$= t ^{\circ}\text{C} = T ^{\circ}\text{K};$	press. = 1 atm
t	T	Sı	S_r	S_v	S	0.2324,5
127	400	36.025	13.959	0.043	50.03	11.627
227	500	37.152	14.626	0.115	51.89	12.061
327	600	38.066	15.167	0.222	53.46	12.425
427	700	38.835	15.626	0.346	54.81	12.738
527	800	39.501	16.024	0.494	56.02	13.020
627	900	40.088	16.375	0.657	57.12	13.277
727	1000	40.611	16.688	0.831	58.13	13.511
827	1100	41.084	16.972	1.020	59.08	13.731
927	1200	41.516	17.231	1.215	59.96	13.957
Saturat	ed vapor (pres	s. = 26.739 mm-l	Hg)			
27	300	41.278	13.104	0.008	54.39	12.642

Table 55 .- (Continued)

III. M. Trautz and H. Ader.²⁴⁶ The following values (C_{vr}) for the rotational component of the specific heat at constant volume were computed from the spectroscopic data of Mecke and Baumann (1933). They concluded that this component attains its full value at a temperature only slightly above 50 °K. As usual, R is the gas constant.

50.0 °K 6.66 10.0 12.5 20.0 25.0 40.0 C_{--}/R 0.1061 0.2252 0.4587 0.5998 0.9359 1.0813 1.3600 1.461

- IV. E. B. Wilson, Jr.,²⁴⁷ has concluded that the distortion caused by the rotation of the molecules affects the thermal properties of the vapor as indicated by the following formulas, ρ being a constant characteristic of the molecule, and a prime indicating what would be the value of the property if the molecule were rigid: $G = G' - \rho RT^2$, $S = S' + 2\rho RT^2$, $C_v = C'_v + C'_v$ $2\rho RT^2$. For water-vapor at very low pressure he gives $\rho = 2.04(10^{-5})$ (°K)-2.
- (H. M. Randall, D. M. Dennison, N. Ginsburg, and L. R. Weber 248 have concluded that no first order computation, such as that of Wilson's, is at all satisfactory when the rotation is so rapid as to cause a displacement of the spectral line in excess of about $\Delta \nu = 5$ cm⁻¹.)

Table 56.—Joule-Thomson Coefficient for Dilated Water-vapor

The Joule-Thomson coefficient (μ) is the decrease in temperature per unit drop in pressure when the expansion is adiabatic. It measures the internal latent heat of expansion. For the more permanent gases, it has been found that the ratio of the decrease in temperature to the associated decrease in pressure "is, for small pressures at all events, constant and independent of both the fall in pressure and the absolute value of the pressure." ²⁴⁹ That is, μ is independent of the pressure. This is nearly, but not quite true for water-vapor. For that, the variation of μ with the pressure, temperature being constant, is small, but not linear in the pressure; and similarly, when the pressure is constant μ is not linear in the temperature. A. Griessmann 250 erred in concluding that μc_p for water-vapor is independent of the temperature, the pressure being constant; the values he accepted for c_n are unsatisfactory. For a discussion of the experimental

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<sup>241</sup> Gordon, A. R., and Barnes, C., J. Phys'l Chem., 36, 1143-1151 (1932).
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²⁴² Mecke, R., Z. Physik, 81, 313-331 (1933).

²⁴⁸ Baumann, W., and Mecke, R., Idem, 81, 445-464 (1933).

²⁴⁴ Freudenberg, K., and Mecke, R., Idem, 81, 465-481 (1933).

²⁴⁵ von Steinwehr, H., Z. Physik, 3, 466-476 (1920).

²⁴⁰ Trautz, M., and Ader, H., Z. Physik, 89, 12-14 (1934).

²⁴⁷ Wilson, E. B., Jr., J. Chem'l Phys., 4, 526-528 (1936).

²⁴⁶ Randall, H. M., Dennison, D. M., Ginsburg, N., and Weber, L. R., Phys. Rev. (2), 52, 160-174 (1937).

³⁴⁰ Buckingham, E., Bull. Bur. Std., 3, 237-293 (S57) (1907).

Table 56.—(Continued)

evidence for the validity of the assumption, often made, that μ obeys the law of corresponding states, see H. N. Davis.²⁵¹

Unit of $P = 1 \text{ kg*/cm}^2 = 0.9678 \text{ atm}$; of $\mu = 1 ^{\circ}\text{C}$ per (1 kg*/cm). Temp. = $t ^{\circ}\text{C}$

I. DK = observations by Davis and Kleinschmidt ^a; KSG' = values computed by F. G. Keyes, L. B. Smith, and H. T. Gerry ²⁵²; KSG" = later values computed by them. ¹⁹⁷ In each case the computation is based on an empirical formula set up by KSG to represent their determinations of the specific volume of dilated water-vapor. The two formulas differ: KSG" supersedes KSG'.

_	DK	KSG'	KSG"	_ P	DK	KSG'	KSG"
P		125°C		_ F		260°C	
1.125	4.801	4.564	4.908	1.60	1.485	1.553	1.495
		145 °C		3.16	1.536	1.557	1.506
1.405	3.730	3.805	3.928	10.55	1.549	1.564	1.539
2.81	3.998	3.816	4.009	14.77	1.548	1.563	1.548
		166 °C		15.11	1.577	1.563	1.549
1.60	2.973	3.159	3.180	20.00	1.551	1.558	1.550
1.76	3.092	3.161	3.182	25.30	1.545	1.550	1.543
2.85	3.209	3.167	3.228	32.40	1.533	1.536	1.526
5.62	3.264	3.173	3.270	39.60	1.511	1.518	1.499
		196 °C				300 °C	
1.60	2.368	2.472	2.405	1.70	1.163	1.202	1.168
3.52	2.409	2.481	2.451	3.16	1.193	1.204	1.174
7.04	2.557	2.485	2.500	8.44	1.192	1.209	1.188
7.04	2.474	2.485		14.00	1.207	1.211	1.199
7.60	2.522	2.484	2.505	15.00	1.197	1.211	1.199
10.55	2.570	2.480	2.500	20,25	1.201	1.210	1.202
10.55	2.490	2.480	2.500	32.70	1.187	1.201	1.199
		225 °C				347 °C	
1.60	1.882	1.985	1.910	1.60	0.932	0.914	0.905
3.16	1.995	1.990	1.930	7.38	0.929	0.918	0.918
7.04	1.978	1.997	1.970	15.00	0.928	0.920	0.922
10.55	1.948	1.995	1.985	35.00	0.919	0.917	0,926
14.77	1.987	1.990	1.990				
20.25	1.953	1.977	1.970				
20.25	2.001	1.977	1.970				

II. Some of preceding DK observations rearranged.

$P \rightarrow I$	1.60	3.16	7.04	10.55	14.77	15.00	20.25	39.60
166	2.973		2 54.6	0 #20	,			
196 225	2.368 1.882	1.995	2.516 1.978	2.530 1.948	1.987		1.977	
260	1.485	1.536	1.770	1.549	1.548		•	1.511
300	0.020	1.193				1.197	1.201	
347	0.932					0.928		

²⁵⁰ Griessmann, A., Z. Ver. deut. Ing., 47, 1852-1857, 1880-1884 (1903).

²⁵¹ Davis, H. N., Proc. Am. Acad. Arts Sci., 45, 241-264 (1910).

²⁸² Keyes, F. G., Smith, L. B., and Gerry, H. T., Mech. Eng., 56, 87-92 (1934).

Table 56.—(Continued)

III. Near state of saturation, A. Griessmann. 250 Values of μ are uncertain by 2 in the first decimal place.

$P \rightarrow t_{\mathtt{sat}} \rightarrow 1$	1.5	119.	.0	126	.5 .8	3. 132.	0
i i	μ	1	μ	t	μ		μ
136.6	4.5	134.1	4.4	136.3	4.2	138.4	4.1
138.0	4.4	135.2	4.4	137.4	4.2	139.4	4.0
141.9	4.3	138.7	4.4	140.9	4.1	143.0	4.1
145.8	4.2	140.2	4.3	142.3	4.1	144.3	4.0
148.6	4.0	144.0	4.3	146.1	4.2	148.1	4.0
150.3	4.0	147.9	4.1	149.9	3.9	151.8	3.8
152.8	3.8	150.7	3.9	152.6	3.9	154.5	3.7
153.7	3.8	152.3	3.9	154.2	3.7	156.0	3.5
133.7	3.0	154.7	3.7	156.5	3.6	158.3	3.6
		155.6	3.7	157.4	3.6	159.2	3.5
		155.0	3.1	137.4	3.0	137,2	0,5
P→	1 5	4.	.0	4	.5	5.	.0
tant-	110.8	142.	9	147	.2	151.	1
	μ	1	μ	1	μ		μ
141.5	3.9	146.9	3.8	150.0	3.6	155.3	3.1
145.0	4.1	148.2	3.8	153.6	3.4	158.7	3.1
146.3	3.9	151.9	3.6	157.1	3.3		
150.1	3.7	155.4	3.4	159.7	3.3	$P \rightarrow$	5.5
153.6	3.6	158.0	3.4			$t_{\text{sat}} \longrightarrow 1$	154.7
156.3	3.5	159.4	3.4			156.8	3.0
157.7	3.4					160.2	3.0
160.0	3.4						
160.9	3.4						
	V. A						

IV. J. R. Roebuck, 253 from H. N. Davis, 251 except as noted.

THERMAL CONDUCTIVITY OF DILATED WATER-VAPOR

In 1931, M. Jakob 258 stated that the only data available for the thermal conductivity of water-vapor are those given by E. Moser 259 for 46 to

Apparently no final account of this work has been published, but the DK values Apparently no final account of this work has been published, but the DK values here given and included in the KSG paper were published by H. N. Davis and J. H. Keenan. They stated (p. 926, see also second paper, p. 253): "The Harvard data have not yet been published, but the definitive experimental results are given in Table 14." Those are the DK values here given. Preliminary reports: H. N. Davis, and R. V. Kleinschmidt. In the last it is stated that the final report of the work has been presented to the Steam Research Committee, and that "we hope to be able to publish it in the near future." But it had not been published in 1929, and apparently it has not yet appeared. Although the values include the third place of decimals, duplicates differ by several units (5 to 8) in the second place.

^b Determination by H. M. Trueblood, ²⁶⁷ $P = 3.86 \text{ kg/cm}^8$.

²⁵⁸ Roebuck, J. R., Int. Crit. Tables, 5, 146 (1929).

²⁰⁴ Davis, H. N., and Keenan, J. H., Mech. Eng., 51, 921-931 (1929); Proc. World Eng. Cong. (Tokyo), 1929, 4, 239-264 (1931).

²⁶⁸ Davis, H. N., Mech. Eng., 46, 85-87, 108 (1924).

²⁵⁰ Kleinschmidt, R. V., Idem, 45, 165-167 (1923); 46, 84-85 (1924); 48, 155-157 (1926).

²⁵⁷ Trueblood, H. M., Proc. Am. Acad. Arts Sci., 52, 731-804 (1917).

100 °C. They are limited to low pressures, and are not very exact. Jakob quotes them as k=45.8 microcal/cm·sec·°C at 46 °C and 56.6 at 100 °C, which are equivalent to 1.92 and 2.37 kiloerg/cm·sec·°C, respectively. From the same source, T. H. Laby and Edith A. Nelson ²⁶⁰ derive the values 1.80 and 2.17 kiloerg/cm·sec·°C. The difference seems to arise from the

Table 57.—Thermal Conductivity of Dilated Water-vapor

S. W. Milverton 262 has reported that his observations between 70 and 95 °C, at pressures between 100 mm-Hg and near-saturation, can be expressed within 1/3 per cent by a formula equivalent to the following, the unit of p being 1 mm-Hg:

 $10^4k = 1.5058 + 0.009081t + 0.001266p - 0.00001130pt \text{ watt/cm} \cdot ^{\circ}\text{C}$

This formula is not to be assumed valid outside the ranges specified, nor for p < 100 mm-Hg; in particular, it leads to a negative temperature coefficient when p exceeds about 1.05 atm. Values defined by it are given in Section I, those lying beyond the limits set by Milverton are enclosed in parentheses.

Unit of p=1 mm-Hg; of P=1 kg*/cm²=735.6 mm-Hg; of $k=10^{-4}$ watt/cm·°C. Temp=l°C

I. Computed by means of Milverton's formula:

$t \xrightarrow{p} t$	100 51.6	200 66.4	300 75.9	400 83.0	500 88.7	75 0 99 .6
55 60	(2.070) (2.110)					
70	2.189	2.236	2.244			
80	2.268	2.305	2.341	0.402	0.440	
90 95	2.348 2.388	2.373 2.407	2.398 2.426	2.423 2.446	$2.448 \\ 2.465$	
100	(2.428)	(2.441)	(2.455)	(2.468)	(2.482)	(2.515)

II. Computed by means of the formula $k = 1.25\eta c_v$ (see text).

$_{P}^{t\rightarrow}$	110	120	130	140	150	160 k	170	180	190	200
1	2.44	2.50	2.56	2.62	2.68	2.74	2.81	2.88	2.95	3.03
2			2.66	2.71	2.76	2.82	2.87	2.93	3.00	3.07
4					2.96	3.00	3.05	3.09	3.13	3.18

III. D. L. Timrot and N. B. Varhaftik, ²⁶³ and N. B. Varhaftik ²⁶⁴ have reported that $k/\eta c_v = 1.361$, as mean value over the range t = 70 to 250 °C and p = 5 to 100 mm-Hg; $k/\eta c_v = 1.416$ at 288.8 °C, and 1.546 at 476.7 °C. Varhaftik states that the temperature coefficient of this ratio between 69 and 476 °C is 3.7 per cent of the value of the ratio at 100 °C.

fact that the values given in the thesis assume for the conductivity of air a value higher than that (2.23 kiloerg/cm·sec·°C) which Laby and Nelson

regard as the best available. For very recent determinations below 100 °C, see Table 57.

Jakob advises that the conductivity at other temperatures and pressures be computed by means of the relation $k = 1.25\eta c_v$, where η is the viscosity and c_v is the specific heat at constant volume. He estimates that the values so computed are uncertain by \pm 5 per cent. The values in Section II of Table 57 have been so computed from the data in Tables 45 and 22.

The thermal conductivity of a mixture of air and water-vapor is not given by the simple additive relation. It is greatest for a mixture containing about 20 per cent of H_2O by volume. Grüss and Schmick ²⁶¹ give the following values, k_a being the conductivity of dry air, t = 80 °C:

$$\%H_2O$$
 7.2 15.0 17.1 19.7 22.5 25.0 30.6 31.2 44.4 51.9 $1000(k-k_a)/k_a$ 20 35 37 36 35 37 26 30 -1 -26

17. REFRACTIVITY OF DILATED WATER-VAPOR

The data in this section are restricted to the optical spectrum; for values of the dielectric constant, see Section 22.

The determinations of the refractivity of water-vapor by C. and M. Cuthbertson ²⁶⁵ were accepted by J. J. Fox and F. G. H. Tate, and given in their compilation ²²⁶; here they will be denoted by the symbol CC. Earlier determinations by Mascart ²⁶⁷ and by Lorenz ²⁶⁸ are not discordant with the Cuthbertson values. The only more recent determinations that have come to the attention of the compiler are those by J. Wüst and H. Reindel ²⁶⁹ and by P. Hölemann and H. Goldschmidt ²⁷⁰; they will be denoted by WR and HG, respectively.

All three (CC, HG, and WR) reduced their observations on the assumption that $r \equiv (n-1)10^6$ is directly proportional to the density (d) of the gas, n being the index of refraction; and they expressed their results in terms of the value that r would have, on that assumption, if the density were such that the vapor contained as many formula weights of H_2O per liter as there are of H_2 in a liter of H_2 at 0 °C and 760 mm-Hg. This

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228 Jakoh, M., Engineering (London), 132, 744-746, 800-804 (1931).
220 Moser, E., Thesis, Berlin, 1913.
220 Laby, T. H., and Nelson, Edith A., Int. Crit. Tables, 5, 215 (1929).
221 Grüss, H., and Schmick, H., Wiss. Veröff. Siemens-Konz., 7, 202-224 (1928).
222 Milverton, S. W., Proc. Roy. Soc. (London) (A), 150, 287-308 (1935).
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224 Varhaftik, N. B., Idem, 31, 6958 (1937) ← 1935, No. 12, 20-23 (1935).
225 Cuthbertson, C. and M., Phil. Trans. (A), 213, 1-26 (1913).
226 Fox, J. J., and Tate, F. G. H., Int. Crit. Tables, 7, 8, 11 (1930).
227 Mascart, E. E. N., Compt. rend., 86, 321-323 (1878).
228 Lorenz, L., Ann. d. Physik (Wied.), 11, 70-103 (1880).
229 Wüst, J., and Reindel, H., Z. physik. Chem. (B), 24, 155-176 (1934).
230 Hölemann, P., and Goldschmidt, H., Idem, 24, 199-209 (1934).
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value we shall denote by r_0 . In making the reduction from r to r_0 , each took for the formula weight of H_2 the value $M_H = 2.016$, and the corre-

Table 58.—Refractivity of Dilated Water-vapor

In each case the value for $\lambda = 5460.7 \,\mathrm{A}$ was determined absolutely by a count of fringes, and the values for other λ 's were determined relative to that. For $\lambda = 5460.7 \,\mathrm{A}$, the values found were $r_0 = 252.7 \pm 0.5$ (CC^a), 256.9 ± 1.0 (WR^a), and 252.1 ± 0.3 (HG^a), where $r_0 = 0.8029 \, r/d$ in the case of CC, and $0.8038 \, r/d$ in the case of the other two (see text). Here d g/l is the density and $r = (n-1)10^6$, n being the index of refraction; λ is the wave-length of the light. The Lorentz-Lorenz expression for the molecular refraction is $R = 1000(n^2-1)M/(n^2+2)d$, M = molecular weight (18.016).

The Cuthbertson dispersion formula is $r_0 = 29190/(118.86 - \lambda^{-2})$, and WR's, after reduction by 1.87 per cent in order to bring their values to the basis of the more accurate value of r_0 obtained by HG for $\lambda = 5460.7$ A, is $r_0 = 34168/(138.89 - \lambda^{-2})$, the unit of λ in each case being 1 μ . The computed values here tabulated have been obtained by means of these equations.

Unit of $\lambda = 1$ A = $10^{-4}\mu = 10^{-8}$ cm; of d = 1 g/l; of R = 1 cm³/gfw. Temp. = t °C

Ref.a→	CC	CC	·WR	CC-WR	., С	
λ	Obs.	ro Comp	outed	Comp.	r/d Obse	rved R
4799.9	254.95	254.89	253.94	95	317.52	3.814
5085.8	253.80	253.84	253.06	78	316.09	3.796
5209.1	253.45	253.44	252.70	74	315.65	3.791
5460.7	252,70	252.71	252.09	62	314.72	3.780
5769.5	251.95	251.95	251.44	51	313.79	3.769
5790.5	251.91	251.90	251.40	50	313.74	3.768
6538.5	250.69	250.67	250.22	45	312.22	3.750
6707.8	250.28	250.26	250.00	26	311.70	3.744

Holemann and Goldschmidt¹⁷⁰. $\lambda = 5460.7$

t	150	250	350	500
R	3.763	3.768	3.761	3.767

C. and M. Cuthbertson; 1936.²⁷⁸ See remarks in text, N_0 and N_e = number of bands observed and calculated, respectively; direct count for $\lambda = 5462.23$ only.

λ	5462.23	4359.54	4078.97	4047.68	3342.42	3126.56	3022.37	2968.13
$N_{ m o} \ N_{ m c}$	776.85 776.85					1064.6 ^b 1064.0 ^b		

a References:

CC C. and M. Cuthbertson.²⁶⁵ WR J. Wüst and H. Reindel.²⁶⁹

sponding value for water, $M_{\rm H_2O}=18.016$. But CC assumed that the density of $\rm H_2$ at 0 °C and 760 mm-Hg is 0.089849 g/l, which is equivalent

b Obviously there is some gross error in these N's for $\lambda = 3126.56$ A.

to a specific volume of 22.438 l/gfw; whereas the others assumed the specific volume under those conditions to be 22.415 1/gfw. Hence, for the CC values $r_0 = 0.8029 \ r/d$, the unit of d being 1 g/l; and for the other two, $r_0 =$ $0.8038 \, r/d$. None of them report the values of d that were used; it appears that the temperature was seldom much below 140 °C. Little physical significance should be attached to the particular values assigned by them to the constants in their interpolation formulas.

A. Bramley 271 has reported that the application of an electric field changes the value of n by an amount in excess of that caused by the attendant change in density arising from electrostriction.

The general subject of optical dispersion has been reviewed by S. A. Korff and G. Breit. 272

Since the foregoing was written, C. and M. Cuthbertson 273 have reported two series of new observations. That of 1934 gave for the green mercury line $r_0 = 253.1$ and that of 1935 gave 252.5; the mean of these two absolute determinations "is almost exactly" 252.7, the value reported in 1913. In reducing their observations they used the same values of the molecular weights as before, but for the density of H2 at 0 °C and 1 atm they used the value 0.08995 g/l, whereas before they used 0.089849 g/l. In their earlier paper they seem to have used wave-lengths in air; in this, wave-lengths in vacuo. They illustrate the agreement of their recent observations with their previously determined dispersion equation by means of the values given at the bottom of Table 58, No being the value defined by that equation when N for $\lambda = 5462.23$ A is 776.85.

P. Hölemann 274 has considered the change in the refraction when a substance passes from the vapor to the liquid state.

18. ABSORPTION OF RADIATION BY WATER-VAPOR

As the spectrum of a gas or vapor consists in large part of numerous narrow lines arranged in bands, and as the observed absorption is the average absorption over a spectral range that is usually greater than the width of a single line and generally great enough to embrace several lines. it is obvious that the simple relation $I = I_0 e^{-kl}$, applicable when the absorption is essentially constant over the spectral range covered by a single observation, will not apply in general to the observed absorption by gases and vapors. For them, the radiation corresponding to the regions between the lines will pass essentially unabsorbed, and in the simplest case the relation will be of the form $I = aI_0 + (1-a)I_0e^{-kl}$ or $(I_0 - I)/I_0 =$ $(1-a)(1-e^{-kl})$, where l is the length along the path of a parallel beam of radiation in the medium between the points where the intensities are I₀

²⁷¹ Bramley, A., J. Franklin Inst., 203, 701-711 (1927).

²⁷² Korff, S. A., and Breit, G., Rev. Mod. Phys., 4, 471-503 (1932). ²⁷³ Cuthbertson, C. and M., Proc. Roy. Soc. (London) (A), 155, 213-217 (1936). ²⁷⁴ Hölemann, P., Z. physik. Chem. (B), 32, 353-368 (1936).

and I, respectively, a is the fraction of the radiation that passes between the lines of absorption, and k is the coefficient of absorption corresponding to the lines in the region observed, and is assumed to be the same for all those lines. If this last assumption is not fulfilled, the expression for the absorption will be more complicated. As l increases, the absorption approaches (1-a), not unity; if l exceeds a certain value, the absorption is essentially independent of l. Furthermore, the apparent coefficient of absorption k', defined by $I = I_0 e^{k'l}$, decreases as l is increased, unless a = 0, in which case k' = k. The decrease is very slow at first but ultimately k' varies as l-1.

Such behavior has been reported. F. Paschen ²⁷⁵ observed that the absorption in the $\lambda=4.3~\mu$ band of CO₂ (pressure = 75 cm-Hg, temp = 17 °C) is essentially as great for a path of 7 cm as for one of 33 cm, that the absorption in the 2.7 μ band of CO₂ is about 28 per cent for a 7-cm path and 43 per cent for a 33-cm path, and that in the water-vapor band near 2.7 μ the absorption is about 60 per cent for the 7-cm path and about 80 per cent for the 33-cm one, the vapor being just under saturation at 100 °C. Whence one obtains for k' the following values, that for the shorter path being given first: CO₂, k'=0.045, 0.017; H₂O, k'=0.13, 0.049 cm⁻¹. In each case, the value for the longer path is markedly less than that for the shorter one.

Even if the absorption varied continuously throughout the spectrum, somewhat similar effects are to be expected in regions in which the variation with λ is great.

One is not justified in assuming that the observed absorption by a gas or vapor under a specified condition follows the exponential law. In general, observations for a single length of path do not suffice for the determination of the absorption for a path of a different length.

The variation of the absorption with the temperature and pressure of the gas or vapor has been studied by E. v. Bahr, 276 continuing work began by K. Angström. 277 She found that many gases, including water-vapor, behave thus for infrared radiation: (1) When the density of the gas is decreased, and the length of the path is correspondingly increased so that the mass of gas traversed per unit cross-section of the path remains unchanged, then the percentage of the incident radiation absorbed is decreased. That is, the apparent coefficient k' decreases more rapidly than the density. (2) If to the expanded gas an inert and transparent gas is added until the total pressure is the same as before expansion, the mass of the expanded gas traversed per unit cross-section of the path remaining unchanged, then the amount of absorption is restored to its value before expansion. This is said to hold for total pressures up to at least 1 atm.

²⁷⁶ Paschen, F., Ann. d. Physik (Wied.), 51, 1-39 (1894). ²⁷⁶ v. Bahr, E., Ann. d. Physik (4), 29, 780-796 (1909) — Diss., Upsala (1909); Idem, 33, 585-²⁷⁷ Angström, K., Ark. Math., Astr., och Fys., 4, no. 30 (1908).

That is, for a given total pressure so produced, the absorption depends solely upon the mass of absorbing gas traversed per unit cross-section of the path, and not at all upon the actual specific volume of that gas. (3) The rate of increase in the absorption with the total pressure, produced as stated, is at first great, but finally becomes zero. For water-vapor she gives the following data for $\lambda = 2.7 \,\mu$, the length of column and the mass of water-

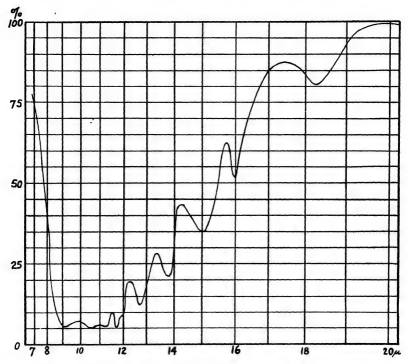


FIGURE 1. Absorption of Infrared Radiation by Water-vapor.

[Adaptation of a figure by H. Rubens and E. Aschkinass, Ann. d. Physik (Wicd.), 64, 584-605 (1898).]

Abscissas = wave-length, unit = $1 \mu = 10^{-4} \text{cm}$; ordinates = per cent of incident radiation that is absorbed by a certain column of water-vapor at a partial pressure of about 1 atm; length of column was about 75 cm. Temperature was somewhat over 100 °C.

vapor remaining unchanged, and the increase in total pressure being produced by adding dry air, which is essentially transparent for that radiation:

Þ	12	100	235	370	570	755	mm-Hg
$100(I_0-I)/I_0$	2.8	4.7	7.2	8.6	10.6	12.1	

(4) Such increase in total pressure does not increase the width of an absorption band, but does increase the intensity of the band at each point, including its maximum. (5) If the pressure is increased solely by heating, the width of an absorption band is increased, but the intensity at the maximum.

mum of the band is unchanged. As regards (4) and (5), water-vapor was not included in her investigations.

That is, the amount of absorption per molecule is influenced by the number of molecular impacts per second, and also by the mean kinetic

Table 59.—Absorption of Radiation by Water-vapor

(See also Figure 1. For absorption of x-rays and γ-rays, see Section 42)

The apparent coefficient of absorption (k') is defined by the relation $I = I_0 e^{-k'l}$; it differs from the true coefficient and depends upon l, unless the strictly monochromatic absorption varies but negligibly over the spectral range for which the individual observations give the average absorption (see text). Furthermore, k' is not proportional to the density of the vapor, and if the vapor is mixed with an inert transparent gas, k' varies with the partial pressure of that gas (see text).

Dreisch (D) and Granath (G) expressed their results in terms of k', believing that the conditions were such that k' is identical with the true coefficient. The other observers expressed theirs in terms of percentage of absorption, Abs = $100(I_0 - I)/I_0$. Granath's data refer to water-vapor saturated at 25 °C (press. = 0.024 atm); all the others refer to a pressure of 1 atm and to pure water-vapor, unless the contrary is indicated. Rubens and Aschkinass (RA) do not give the exact temperature of the vapor they used, stating merely that the tubular container was heated above 100 °C, so as to avoid condensation; the value of the specific volume here used is that corresponding to 100 °C; the pressure was 1 atm.

The values of the absorption reported by L. R. Weber and H. M. Randall ²⁸¹ seem to have been given solely for the purpose of indicating the intensities of the several absorption lines; coefficients of absorption cannot be derived from them and the accompanying data.

Unit of $\lambda = 1 \mu = 10^4 \text{A}$; of Abs = 1%; of $l = 1$ cm	1; of $v/m = 1 \text{ cm}^3/g$; of $k' = 1 \text{ cm}^3$. Temp. = $t ^{\circ}\text{C}$
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λ	Abs	I	v/m	ŧ	1000k'	k'v/m	Ref.a
0.190	18	29	43310	25	7.0	300	G
0.195	51	241	43310	25	3.0	130	G
0.205	24	241	43310	25	1.2	50	G
0.95	8	109	1801	127	0.76	1.4	H
1.12	14	109	1801	127	1.4	2.5	\mathbf{H}
1.35	11.0	25	1677	100	4.6	7.7	D
1.37	20.3	25	1677	100	8.7	14.6	D
1.37	<i>7</i> 5	109	1801	127	12.7	22.9	H
1.404	38.3	25	1677	100	19.3	32.4	D
1.45	24.8	25	1677	100	11.4	19.1	D

²⁷⁸ Paschen, F., Ann. d. Physik (Wied.), 50, 409-443 (1893); 51, 1-39, 40-46 (1894); 52, 209-237 (1894); 55, 287-300 (1894).

²⁷⁰ Schmidt, H., Ann. d. Physik (4), 29, 971-1028 (1909).

²⁶⁰ Fowle, F. E., Astroph. J., 42, 394-411 (1915); Smithsonian Misc. Collect., 68, No. 8 (publ. 2484), (1917); Smithsonian Physical Tables.

²⁸¹ Weber, L. R., and Randall, H. M., Phys. Rev. (2), 40, 835-847 (1932).

Table 59.—(Continued)

		-	able Ja	(Commune	*/		
λ	Abs	I	v/m	*	1000k'	k'v/m	Ref.
1.50 1.80 1.83 1.85 1.885	10.1 8.8 84 37.3 47.6	25 25 109 25 25	1677 1677 1801 1677 1677	100 100 127 100 100	4.2 3.7 16.8 18.7 25.8	7.1 6.2 30.2 31.4 43.3	D D H D
1.9 1.935 1.97 2.0 2.48	74 37.5 25.9 8.6 93	109 25 25 25 25 109	1801 1677 1677 1677 1801	127 100 100 100 127	12.4 18.8 12.0 3.6 24.4	22.3 31.5 20.1 6.0 43.9	H D D H
2.55 2.585 2.618 2.65 2,82	47.5 80.0 91.8 77.2 91	25 25 25 25 109	1677 1677 1677 1677 1801	100 100 100 100 127	25.8 64.4 90.2 59 22.1	43.3 108 151 99 39.8	D D D H
3.19	40	109	1801	127	4.7	8.5	H
3.26	27	109	1801	127	2.9	5.2	H
5.25	85	109	1801	127	17.4	31.3	H
7.0	75	75	1677	100+	18.5	31	RA
7.55	90	104	1801	127	22.1	39.8	H
7.90	72	104	1801	127	12.2	22.0	H
8.0	40	75	1677	100+	6.8	11.4	RA
8.2	32	104	1801	127	3.7	6.7	H
9.0	5	75	1677	100+	0.7	1.1	RA
10.0	7	75	1677	100+	1.0	1.7	RA
11.0 11.5 11.7 12.0 12.4	6 10 5 9 20	75 75 75 75 75	1677 1677 1677 1677 1677	100+ 100+ 100+ 100+ 100+	0.8 1.4 0.7 1.3 3.0	1.3 2.3 1.1 2.2 5.0	RA RA RA RA
12.4	34	104	1801	127	4.0	7.2	H
12.8	13	75	1677	100+	1.9	3.2	RA
13.0	18	75	1677	100+	2.7	4.5	RA
13.3	47	104	1801	127	6.1	11.0	H
13.4	28	75	1677	100+	4.4	7.4	RA
13.9	22	75	1677	100+	3.3	5.6	RA
14.0	28	75	1677	100+	4.4	7.4	RA
14.3	43	75	1677	100+	7.5	12.5	RA
14.4	61	104	1801	127	9.0	16.3	H
15.0	35	75	1677	100+	5.7	9.6	RA
15.5	52	75	1677	100+	9.8	16.4	RA
15.6	76	104	1801	127	13.7	24.7	H
15.7	63	75	1677	100+	13.2	22.2	RA
16.0	52	75	1677	100+	9.8	16.4	RA
16.5	74	75	1677	100+	18.0	30.1	RA
17.0	85	75	1677	100+	25.3	42.4	RA
17.0	88	104	1801	127	20.4	36.7	H
17.0	52	32.4	1801	127	22.6	40.7	H
17.3	61	32.4	1801	127	29.0	52.2	H
17.5	88	75	1677	100+	28.3	47.5	RA
18.0	82	75	1677	100+	22.9	38.4	RA
18.3	80	75	1677	100+	21.5	36.1	RA
18.3	61	32.4	1801	127	29.0	52.1	H
18.5	83	75	1677	100+	23.6	39.6	RA
19.0	93	75	1677	100+	35.5	59.5	RA

A	/	~	•
Table	59	Continued)

λ	Abs	I	v/m	ŧ	1000k'	k'v/m	Ref.
19.2	82	32.4	1801	127	52.9	95.2	H
19.5	98	<i>7</i> 5	1677	100 +	52.1	87.4	RA
19.8	84	32.4	1801	127	56.5	101.8	H
20,0	99	75	1677	100 +	61.3	103	$\mathbf{R}\mathbf{A}$
20.3	81	32.4	1801	127	51.2	92.2	H
32	60.4	40	1723	110	23.2	39.9	RWg
52	99.3	40	1723	110	124	213	RWg
108	80.4	40	1905	(?)	40.8	78	RWd
110	80.4	40	1723	110	40.8	70.3	RWg
314	50.8	40	1723	110	17.7	30.5	RWg

Positions of other maxima (m) and of regions (w) of very little absorption.

λ	Ref.ª	λ	Ref.a
47 w	We	79 m	We
50 m	We, Wi	79.3 m	Wi
52.5 m	Wi	90.9 m	W_i
54 w	We	91 w	We
58 m(?)	We	103 m (?)	We
56.6 m	Wi	108.9 m	W_i
62 w	We	115 w	We
63.7 m	Wi	116.8 m	Wi
66 m	We	125 w (?)	We
69.6 m	Wi	131.8 m	W_i
74.5 m	Wi	138 w (?)	We
75 w	We	167 m	W_i

Diathermacy of moist air for the complete radiation from an enclosure at the temperature t °C. Length of tube = 250 cm. Pressure = 1 atm; temp. = 70 °F = 21.1 °C, moisture content = 0.032 mm of precipitable water, giving lm/v = 0.0032 g/cm², v/m = 78 1/g of vapor, and relative humidity = 70 per cent. (In the legend to the graphs, the temperature of the moist air is incorrectly given as 70 °C.) ²⁵²

t	510	590	735	760	780°	820	850
$100I/I_{0}$	91.3	92.2	92.6	92.6	92.6	92.7	92.9
1000k'	0.36	0.33	0.31	0.31	0.31	0.30	0.29
k'v/m	2.6	2.4	2.2	2.2	2.2	2.2	2.1

[&]quot; References and remarks:

D T. Dreisch, 258 quoted by J. Becquerel and J. Rossignol. 254 The significance of the values tabulated by Dreisch is not entirely clear. The heading of the column indicates that the values are k' in our notation, and it is stated that l=1 meter; but it seems that he means by the last merely that the unit of k' is $1 \, \text{m}^{-1}$, which is the interpretation adopted by Becquerel and Rossignol, and used in this compilation. Neither the temperature, the pressure, nor the density of the vapor is explicitly stated, but the text indicates that the vapor was probably saturated at $100\,^{\circ}\text{C}$, corresponding to a specific volume of $1677\,\text{cm}^{\circ}/\text{g}$. Assuming this and the preceding interpretation of the tabular data to be correct, one finds that the ordinates of his Fig. 2 should each be 0.3 of the corresponding tabular value, which they are. The values here assigned to him have been derived from his data on the basis of

²⁸² Brown, S. L., Phys. Rev. (2), 21, 103-106 (1923).

²⁸³ Dreisch, T., Z. Physik, 30, 200-216 (1924).

²⁸⁴ Becquerel, J., and Rossignol, J., Int. Crit. Tables, 5, 269 (1929).

Table 59.—(Continued)

these assumptions. Becquerel and Rossignol appear to have erred in stating that the values of k' refer to vapor at 0 °C and 1 atm.

- G L. P. Granath 285 used vapor saturated at 25 °C.
- G. Hettner 286 used vapor at 127 °C and 1 atm. H
- H. Rubens and E. Aschkinass, 267 vapor saturated at 100 °C was passed RA into the middle of a metal tube open at each end and heated somewhat above 100 °C.
- H. Rubens and R. W. Wood. 288 RWd
- H. Rubens and H. v. Wartenberg 289 used vapor at 110 °C and 1 atm, RWg tube open at ends.
- We W. Weniger.290
- H. Witte. 201 Wi
- b Neither the temperature nor the pressure is stated.
- ° Using a column of moist air 51 cm long, v/m = 100500 cm³/g of vapor, t = 800 °C, W. W. Coblents we found absorption = 0.9 per cent, which corresponds to 1000k' = 0.18, k'v/m = 18. For a column of dry air of the same length, the absorption was about 0.09 per cent, "which is the magnitude of the errors of observation." The pressure was 1 atm in each case. The measurement was incidental to another investigation.

energy of the impacting molecules; but the intimate effects of these two influences seem to differ.

In general, determinations made at a single partial pressure, a single temperature, and a single wave-length do not suffice for the determination of the absorption under other conditions.

Observations by F. Paschen ²⁷⁸ indicate that the infrared radiation from heated gases is of thermal origin, and satisfies Kirchhoff's law, cases involving obvious chemical and electrical effects being excluded; but the evidence is not entirely convincing.²⁷⁹ See also, C. Schaefer and F. Matossi, "Das Ultrarote Spektrum," p. 104, 1930.

Tables for use in the determination of the effect of moisture upon the atmospheric transmission of solar and of terrestrial radiation have been published by F. E. Fowle, 280 together with his observations bearing thereon.

19. Emissivity of Water-vapor

The only direct measurements of the emissivity of water-vapor and of its variation with the thickness of the layer of vapor seem to be those mentioned in Table 60. Earlier estimates based upon the values of the spectral absorptivity in the regions effective—the bands at $\lambda = 2.67 \,\mu$ and $6.6 \,\mu$.

- 288 Granath, L. P., Phys. Rev. (2), 34, 1045-1048 (1929) → 33, 1073 (A) (1929).
- ²⁶⁰ Hettner, G., Ann. d. Physik (4), 55, 476-496 (1918). ²⁶⁷ Rubens, H., and Aschkinass, E., Ann. d. Physik (Wied.), 64, 584-605 (1898) = Astrophys. J., 8, 176-192 (1898).
 - 288 Rubens, H., and Wood, R. W., Verh. deut. physik. Ges., 13, 88-100 (1911).
- 250 Rubens, H., and v. Wartenberg, H., Physik. Z., 12, 1080-1084 (1911) = Verh. deut. physik. Ges., 13, 796-804 (1911).
 - 200 Weniger, W., J. Opt. Soc. Amer., 7, 517-527 (1923).
- ²⁰¹ Witte, H., Z. Physik, 28, 236-248 (1924). ²⁰² Coblentz, W. W., Proc. Nat. Acad. Sci., 3, 504-505 (1917); also Sci. Papers Bur. Stand., 15, 529-535 (S357) (1920).

Table 60.—Emissivity of Water-vapor

 ϵ = ratio of the net radiation from the layer of water-vapor, or of a mixture of the vapor and CO₂-free air, to that from the ideal radiator (black body) at the same temperature. The intensity of radiation from the ideal radiator may be found in Table 288; τ = thickness of the layer of radiating gas; p = partial pressure of the vapor. Temp. = t °C = t_F °F.

Unit of $\tau = 1$ cm; of $\phi = 1$ atm; ϵ is dimensionless

I. Pure vapor. E. Schmidt 207 ; values confirmed by E. Eckert. 295 The following values have been read from Schmidt's graph; p=1 atm in all cases.

au ightarrow t	0.96	2.00	3.02	4.02 —— 100ε —	6.00	12.0	18.2
100	9.0	14.4 .	17.9	20.8	23.6	30.2	34.3
200	8.1	13.2	16.6	19.4	22.3	29.0	33.2
300	7.2	11.8	15.2	17.8	20.8	27.6	31.8
400	6.1	10.2	13.7	16.0	19.2	26.0	30.2
500	5.2	8.8	12.0	14.4	17.4	24.3	28.4
600	4.4	7.8	10.5	12.6	15.6	22.6	26.6
700	3.8	6.7	9.2	11.2	13.8	20.7	24.8
800	3.2 .	6.0	8.2	10.0	12.4	19.0	23.0
900	2.8	5.2	7.3	9.0	11.3	17.4	21.4
1000	2.6	4.8	6.6	8.1	10.3	16.0	20.0

II. Vapor mixed with air. H. C. Hottel and H. G. Mangelsdorf.²⁹⁸ The following values have been read from their graph. In all cases, $\tau = 51.2$ cm (1.68 ft.) and total pressure = 1 atm.

	¢ τ→ ¢→	0.256 0.00 50	0.51 0.010	1.02 0.020	2.04 0.040	4.09 0.080	8.5 0.167	25.6 0.50	51.2 1.00
tr	t				100ε				
200	93	1.36	2.6	5.0	8.5	14	23	39	50
400	204	1.15	2.2	4.1	7.6	12.5	21	36	48
600	316	0.98	1.9	3.6	6.7	11.0	18.5	34	45
800	427	0.82	1.6	3.2	5.9	10.0	16.8	31	43
1000	538	0.71	1.4	2.8	5.2	9.0	15.2	29	41
1200	649	0.63	1.3	2.5	4.6	8.0	14.0	27	39
1400	760	0.57	1.10	2.2	4.1	7.2	12.5	26	37
160 0	871		0.96	1.98	3.7	6.5	11.5	24	35
1800	982		0.88	1.72	3.2	5.8	10.2	22	34
2000	1093		0.75	1.51	2.9	5.3	9.4	21	32
2200	1204		0.70	1.38	2.6	4.8	8.6	19	30
2400	1316		0.61	1.20	2.4	4.4	8.0	18	29
2600	1427		0.52	1.08	2.1	4.0	7.2	17	27
2800	1538			0.98	1.96	3.6	6.7	16	26
3000	1649			0.89	1.80	3.3	6.2	15	25

²⁸³ Schack, A., Z. Ver. deut. Ing., 68, 1017-1020 (1924); Z. techn. Physik, 5, 267-278 (1924); Idem, 7, 556-563 (1926).

and the region $\lambda=12$ to 25 μ —had been made by A. Schack.²⁹⁸ They involve a number of simplifying assumptions, including the following:

- 1. Where the absorption is the greatest in any band, the observed relative amount of radiation transmitted is exponentially related to the thickness; $I = I_0 e^{-k_m x}$, k_m being independent of x. In general, this is not true (see Section 18).
- 2. The integral effect of any given band can be represented by an expression of the form

$$I_0 \int_{\lambda_0}^{\lambda_0 + \Delta \lambda} e^{-k_m x (\lambda - \lambda_0)/\Delta \lambda} d\lambda = \frac{I_0 \Delta \lambda (1 - e^{-k_m x})}{k_m x}$$

the band extending from λ_0 to $\lambda_0 + \Delta \lambda$.

3. The value of k_m observed when the temperature and pressure are relatively low will apply when they are high. Actually, the individual lines are broadened and the local variations in intensity throughout the band are, in part, wiped out as the temperature and the pressure are increased, either singly or together.

Owing to such assumptions and to the fact that the value found for k_m depends upon experimental details, these estimates are unsatisfactory, although they were of great value when made, being the only estimates then available.

M. Jakob ²⁵⁸ has discussed them, and has compared them with the results obtained by Schmidt; and H. C. Hottel ²⁹⁴ has published an English paraphrase of Schack's work, extended by graphs and some elaboration of detail.

The intensity of the radiation emitted at a given temperature by a given volume of water-vapor when mixed with a nonabsorbing and nonradiating gas is not determined solely by the amount of the vapor contained in that volume, *i.e.*, Beer's law is not valid for such a mixture.^{295, 296}

20. LUMINESCENCE OF WATER-VAPOR

The types of luminescence here considered are these: Fluorescence, including phosphorescence; the Rayleigh scattering (also called the Tyndall effect); the Raman scattering; and the scattering of x-rays. The distinctive characteristics of these several effects, except the last, are considered in Section 39.

²⁰⁴ Hottel, H. C., Trans. Am. Inst. Chem. Eng., 19, 173-205 (1927); J. Ind. Eng. Chem., 19, 888-894 (1927).

²⁰⁰⁶ Eckert, E., Forschungsheft, 387, 1-20 (1937).

²⁰⁰ Schmidt, E., and Eckert, E., Forsch. Gebiete Ingenieurw., 8, 87-90 (1937).

²⁹⁷ Schmidt, E., Forsch. Gebiete Ingenieurw., 3, 57-70 (1932).

²⁰⁰⁸ Hottel, H. C., and Mangelsdorf, H. G., Trans. Am. Inst. Chem. Eng., 31, 517-549 (1935).

Fluorescence.

When water-vapor at atmospheric pressure is illuminated by light from the mercury line $\lambda=2537\mathrm{A}$, it emits an intense ultraviolet fluorescence having a continuous spectrum which has a maximum near 2537A, and extends several thousands of wave-numbers (cm⁻¹) toward the visible spectrum; its intensity is some thousands of times greater than that of the Raman-scattered light.²⁰⁹

When moist nitrogen is exposed to radiation of high frequency (ultra-Schumann), it luminesces, and the spectrum of the luminescent light contains the so-called water-band. This was described as a fluorescence of water-vapor. But it is now agreed that the "water-band" actually arises from the OH molecule. Consequently, the observed luminescence cannot properly be described as a fluorescence of water-vapor. G. H. Dieke ³⁰¹ so remarks, and explains the dissociation as arising from "impacts of the second kind" between H₂O molecules and excited N₂ molecules; in which case, the band should vanish if the N₂ were eliminated.

H. Neuimin and A. Terenin 302 have stated that radiation from the Schumann region excites luminescence in water-vapor as a result of photo-dissociation combined with excitation of the resulting OH radical. The hydroxyl band at $\lambda=3062\mathrm{A}$ is particularly strong. The luminescence is strongest when the pressure of the vapor is about 0.8 mm-Hg; it is reduced by the presence of CO or of $\mathrm{H_2}$, but not by that of $\mathrm{N_2}$ or of argon.

In speaking of the reaction that occurs in a mixture of H_2 and O_2 , S. Horiba 303 has stated that in the transition region from nonexplosive to explosive reaction at the lower critical pressure there is luminescence, although the reaction velocity is measurably small, and heating seldom occurs.

Rayleigh Scattering.

The depolarization (see Section 39) of the light scattered by water-vapor is $\rho = 0.0199$, the density of the gas being low.³⁰⁴ Such a defect in polarization has been explained by Lord Rayleigh ³⁰⁵ and by M. Born ³⁰⁶ as being due to an anisotropy of the molecule; but I. R. Rao ³⁰⁷ has concluded that it is the anisotropy of the oxygen ion that is involved. For a discussion of the anisotropy of the H₂O molecule and of its ions, see Section 9.

No direct measurement of the ratio of the intensity of the light laterally scattered by water-vapor to that of the incident light has been found, but from other ratios obtained by others, W. H. Martin and S. Lehrman ³⁰⁸ have computed the value $Ir^2/EV = 1.06 \times 10^{-8}$ per atmosphere, as applying to water-vapor at 27 °C. Here I = intensity of the scattered light at the distance r from the scattering vapor of volume V, and E is the intensity of the incident (exciting) light.

²⁰⁰ Rasetti, F., Nuovo Cim., N. S., 8, 191-193 (1931).

Raman Scattering.

As each frequency in the incident radiation gives rise to its own series of Raman lines, the interpretation of the observed spectrum is difficult, or even ambiguous, unless the incident radiation is unifrequent, which it usually is not. For a discussion of the general subject and the earlier data see K. W. F. Kohlrausch, "Der Smekal-Raman Effekt," 1931.

Table 61.—Raman Spectrum of Water-vapor

 $\delta \nu =$ difference between the wave-number of the Raman line and that of the associated line in the incident radiation; $\lambda_R = \delta \nu$ is the wave-length corresponding to the fundamental frequency responsible for $\delta \nu$, and presumably has a representative of nearly the same value in the infrared spectrum of the vapor; No. = number of components observed.

No.		0	ν		Ref.
1 3		3655 3654	1648	984	DK JW
2 (?) 1 1	3804 (?)	3650 3655 3654.5	none	none	Rk Ro Be
$\lambda_R \longrightarrow$	2.63	3646 ^b 2.74	6.06	10,2	Uk

Unit of $\delta \nu = 1 \text{ cm}^{-1}$; of $\lambda_R = 1 \mu = 10^4 \text{A} = 10^{-4} \text{ cm}$

* References and remarks:

- Be D. Bender. 800
- DK P. Daure and A. Kastler all used vapor saturated at 130 °C; found a single, fine Raman line with no satellite of comparable intensity.
- JW H. L. Johnston and M. K. Walker. 311
- Rk D. H. Rank ³¹² used vapor at atmospheric pressure; sought for the JW lines at $\delta \nu$ 1648 and 984, but failed to find them. On long exposure found a line ($\nu = 21950$) that JW assigned to a 984 cm⁻¹ shift from $\lambda = 4358$ A, but which can just as well be regarded as a 3650 cm⁻¹ shift from $\lambda = 3906$ A. The $\delta \nu = 3804$ cm⁻¹ is quite doubtful; the line was very weak, though the exposure was for 93 hours.
- Ro I. R. Rao as concludes that the data available early in 1934 indicate that there is but a single line, that for which $\delta v = 3655$ cm⁻¹.
- Uk S. A. Ukholin. 314
- ^b This is for density $\rho=0.055$ g/cm³; at low density the line becomes double, $\delta\nu=3639$ and 3653 cm⁻¹.

³⁰⁰ Wood, R. W., Phil. Mag. (6), 20, 707-712 (1910); Wood, R. W., and Hemsalech, G. A., Idem, 27, 899-908 (1914); Meyer, C. F., and Wood, R. W., Idem, 30, 449-459 (1915).

⁸⁰¹ Dieke, G. H., Proc. Akad. Wet. Amsterdam, 28, 174-181 (1925).

³⁰² Neuimin, H., and Terenin, A., Acta Physicochim. URSS, 5, 465-490 (1936).

³⁰⁸ Horiba, S., Sci. Rep. Tôhoku Imp. Univ. (Sendai), (1) Honda Anniv. Vol., 430-443 (1936).

⁸⁰⁴ Rao, I. R., Indian J. Phys., 2, 61-96 (1927).

⁸⁰⁶ Lord Rayleigh, Phil. Mag. (6), 35, 373-381 (1918).

⁸⁰⁰ Born, M., Verh. deut. physik. Ges., 19, 243-264 (1917); 20, 16-32 (1918).

⁸⁰⁷ Rao, I. R., Indian J. Phys., 2, 435-465 (1928).

⁸⁰⁸ Martin, W. H., and Lehrman, S., J. Phys'l Chem., 26, 75-88 (1922).

⁸⁰⁰ Bender, D., Phys. Rev. (2), 47, 252 (L) (1935).

⁸¹⁰ Daure, P., and Kastler, A., Compt. rend., 192, 1721-1723 (1931).

Scattering of X-rays.

H. Gajewski ⁸¹⁵ has found that his observed intensity (I) of the x-rays scattered by water-vapor at an angle θ to the direction of the incident beam agrees within experimental error with that calculated for the triangular molecule with the O-H distance = 0.86A and the H-H = 1.28A, which distances (see Table 16) are those published by R. Mecke. ³¹⁶ Account was taken of the effect of the hydrogen as well as of the oxygen, of the interaction between the two, and of the Compton effect. His values are as follows, the unit of intensity being that at $\theta = 0$:

21. SPECTRA OF WATER-VAPOR

Absorption Spectrum.

The absorption spectrum of water-vapor extends from the far infrared to the far ultraviolet (see Table 62), and consists of a series of bands, each composed of numerous lines. In places, several bands overlap. Only recently has it been possible to arrange the bands in a satisfactory order, and to analyze their structure. Among the earlier attempts to analyze the bands of longer wave-length, may be mentioned those of C. R. Bailey, ³¹⁷ C. R. Bailey, A. B. D. Cassic and W. R. Angus, ³¹⁸ H. Deslandres, ³¹⁹ J. W. Ellis, ³²⁰ A. Eucken, ³²¹ G. Hettner, ³²² F. Hund, ³²³ P. Lueg and K. Hedfeld, ³²⁴ R. Mecke, ³¹⁶ E. K. Plyler, ³²⁵ W. W. Sleator and E. R. Phelps, ³²⁶ and H. Witt, ³²⁷

The papers from Mecke's laboratory in 1932 328 should be consulted for procedure and comments, but the numerical analysis then published is superseded by that of 1933.329 The result of this analysis is given in Tables 64

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41 Johnston, H. L., and Walker, M. K., Phys. Rev. (2), 39, 535 (L) (1932).
   812 Rank, D. H., J. Chem'l Phys., 1, 504-506 (1933).
   818 Rao, I. R., Phil. Mag. (7), 17, 1113-1134 (1934).
   814 Ukholin, S. A., Compt. rend. Acad. Sci. URSS, 16, 395-398 (1937).
   815 Gajewski, H., Physik. Z., 33, 122-131 (1932).
   816 Mecke, R., Physik. Z., 30, 907-910 (1929).
   an Bailey, C. R., Trans. Faraday Soc., 26, 203-212, 213-215 (1930).
   818 Bailey, C. R., Cassie, A. B. D., and Angus, W. R., Idem, 26, 197-202 (1930).
   219 Deslandres, H., Compt. rend., 180, 1454-1460, 1980-1986 (1925).
   820 Ellis, J. W., Phil. Mag. (7), 3, 618-621 (1927).
<sup>321</sup> Eucken, A., Verh. deut. physik. Ges., 15, 1159-1162 (1913); Jahrb. d. Radio-akt., 16, 361-411 (1920); Z. Elektroch., 26, 377-383 (1920).
   823 Hettner, G., Z. Physik, 1, 345-354 (1920).
   828 Hund, F., Idem, 43, 805-826 (1927).
   824 Lueg, P., and Hedfeld, K., Idem, 75, 512-520 (1932).
   825 Plyler, E. K., Phys. Rev. (2), 39, 77-82 (1932).
   836 Sleator, W. W., and Phelps, E. R., Astroph. J., 62, 28-48 (1925).
   27 Witt, H., Z. Physik, 28, 249-255 (1924).
<sup>228</sup> Mecke, R., Die Naturwiss., 20, 657 (1932); Z. physik. Chem. (B), 16, 409-420, 421-437 (1932); Mecke, R., and Baumann, W., Physik. Z., 33, 833-835 (1932).

<sup>229</sup> Mecke, R., Z. Physik, 81, 313-331 (1933); Baumann, W., and Mecke, R., Idem, 81, 445-464 (1933); Freudenberg, K., and Mecke, R., Idem, 81, 465-481 (1933).
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Table 62.—Lines and Bands in the Absorption Spectrum of Water-vapor*

Herein are listed the recorded wave-lengths of the maxima or centers of the several bands, of the isolated lines that may not be involved in the fine-structure of the bands, and of certain series of lines (B, Ra, WbR) forming the fine-structure of bands. The last are included solely as illustrations of the nature of such structure; for additional data, reference should be made to the sources listed in Table 66. All wave-lengths recorded by B and by Ra are listed, but only the distinct maxima observed by D and by WbR are given.

Unit of $\lambda = 1 \mu = 10^4 A = 10^{-4} \text{ cm}$; of $\nu = 1 \text{ cm}^{-1}$											
λ	ν	R^a	Ref.b		λ		ν	R^a	Ref.		
400°	25.0	0	Ba		82.4		121.4		K		
250°	40.0		Ba		81.988		121.97		WrR		
170° 167.	58.8		Ba		79.3		126.1		Wi		
167.	59.9		Wi		79.		126.		Ru1, 8		
150.	66.7		Wi		78.63		127.2		WrR		
134.7	74.2		WrR		78.0		128.2		Ru ²		
132.3	<i>7</i> 5.6		WrR		77.66		128.77		WrR		
132.2	75.6		Ru^2		75.6		132.3		K		
131.8	75.9		Wi		75.32		132.77		WrR		
127.8	78.2		WrR		74.5		134.2		Wi		
126.5	79.0		WrR		72.4		138.1		K		
125.6	79.6		WrR		72.2		138.5		Ru*		
121.7	82.2		WrR		71.79		139.30		WrR		
116.8	85.6		Wi		70.98		140.88		WrR		
113.1	88.4		WrR		69.6		143.7		Wi		
111.7	89.5		WrR		66.62		150.10		WrR		
108.9	91.8		Wi		66.0		151.5		K.		
108.1	92.5		WrR		66.		152.		Ru ¹		
105.8	94.5		Ru²		65.8		152.0 153.52		Ru ² WrR		
104.03	96.13		WrR		65.14						
103. (?)	97.1		Ru¹		63.7		157.0		Wi		
100.96	99.05		WrR Wi		63.34 60.01		157.88 166.64		WrR		
99.5 99.415	100.5 100.59		WrR		58.2		171.8		WrR K		
98.559	100.39		WrR		58 (?)		172.		Ru¹		
							173.3		Ru		
95.613	104.59		WrR		57.7 56.6		176.7		Wi		
94.541 94.	105.77 106.		WrR StW		56.3		177.6		K		
93.199	107.30		WrR		52.5		190.5		Wi		
92.662	107.92		WrR		52.0		192.3		K		
90.9	110.0		Wi		52.		192.3		StW		
89.919	111.21		WrR		50.		200.		Wi		
88.520	112.97		WrR		50.		200.		Ru1, 2		
85.662	116.74		WrR		49.8		200.8		K		
85.0	117.6		K		49.0		204.1		Ru²		
84.690	118.08		WrR		48.0		208.3		K		
83.196	120.20		WrR		44.2		226.2		ĸ		
83.	120.20		Wi		44.1		226.8		Ru²		

^{*} Since this table was written the region 18 to 75 μ has been carefully mapped and analyzed by Randall and associates, 38 to 170 μ by Barnes and associates, and the ultraviolet spectrum beyond $\lambda = 1785~\mu$ by Price (see text); and the 6324Å band in the spectrum of the low sun has been studied by V. N. Kondratjev and D. I. Eropkin.

Table 62.—(Continued)

λ	ν	R^a	Ref.b	λ	v	R^a	Ref.b
40.6	246.3		K	12.82	780.0		RuHt
40.0	250.0		Ru²	12.65	790.5	15	WbR
38.6	259.1		K "	12.42	805.2	S	RuHt
35.7	280.1		Ru ²	12.4	806.	-	RuA
35.6	280.9		K	12.14	823.7	7	WbR
33.0	303.	S	Ht	11.89	841.0	_	RuHt
32.9	304. 323.	s	Ru² Ht	11.77 11.66	849.6	7	WbR
31.0	323. 327.	5	Ru ²	11.60	857.7 862.	S	RuHt RuA
30.6 28.9	346.		Ru ²	11.47	871.8		RuHt
26.6	376.		Ru ²	11.24	889.7		RuHt
25.0 25.0	400.		Ru²	11.06	904.2	5	WbR
24.72	404.5	38	WbR	10.94	914.1	S	RuHt
23.81	420.0	66	WbR	10.9	917.	_	RuA
23.8	420.		Ru²	10.80	925.9		RuHt
22.9	437.		Ru ²	10.66	938.1		RuHt
22.61	442.1	31	WbR	10.42	959.7	9	WbR
22.38	446.8	42	WbR	10.30	970.9	S	RuHt
21.81	458.5	65	WbR	9.98	1002.		RuHt
21.6	463.	5 0	Ru ²	9.74	1027.	S	RuHt
21.12	473.5	53 28	WbR WbR	9.50	1053.		RuHt
20.62	485.0 488.	20	Ru ²	9.30 6.296	1075.		RuHt
20.5 20.29	492.8	24	WbR	6.2673	1588.3 1595.6	С	M(T) SPh, BC
19.87	503.3	30	WbR	6.26	1597.	c	Many
19.8	505.	s	Ht	3.168	3156.	c	PIS
19.8	505.	-	RuA	3.168	3156	c	M(T)
19.7	508.		Ru²	3.15	3175		Many
19.3	518.	32	WbR	3.109	3216		SPh
19.2	521.		Ru^2	2.672	3742	С	SPh
19.02	525.8	21	WbR	2.663	3755		M(T), BC
18.37	544.4	4.0	RuHt	2.66	3759		Many
18.34	545.2	19	WbR	2.618	3820	9.0	D
18.21	549.1	14	WbR WbR	2.05	4878		Many
17.57	569.2	16		2.00	5000	0.0	Many
17.5 17.33	571. 577.0		RuA RuHt	1.885 1.875	5305 5333	2.6	D M(T)
17.33	578.	s 14	WbR	1.87	5348		M(T) Many
17.0	588.		Ht	1.870	5348		SPh
16.89	592.1	14	WbR	1.46	6849		Many
16.80	595.2		RuHt	1.45	6896	С	McU
16.53	605.0	14	WbR	1.404	7122	1.9	D
16.00	625.0		RuHt	1.382	7236	С	SPh
15.99	625.4	40	WbR	1.38	7246	c	McU
15.73	635.7	32	WbR	1.379	7252		M(T)
15.7	637		RuA	1.37	7299		Many
15.62 15.17	640.2 659.2	25 25	RuHt WbR	1.16	8621		Many
14.98	667.6	23	RuHt	1.135 1.13	8810 8850		M(Ť) McU
14.5	690.	35	WbR	1.13	8850	С	Many
14.32	698.3	s	RuHt	0.964868	10364.1	2	
14.3	699	3	RuA	0.964559	10367.47	5	B B B
13.62	734.2		RuHt	0.964506	10368.0	5	B
13.50	740.7	17	WbR	0.963758	10376.0	2 5 6 6	В
13.4	746.		RuA	0.963616	10377.6		В
13.34	749.6	S	RuHt	0.958184	10436.4	3	В
13.06	765.7	17	WbR	0.958112	10437.2 \$	4	В

Table 62.—(Continued)

							D 41
λ	v	R^{a}	Ref. b	λ	<i>v</i>	R ^s	Ref. b
0.958005	10438.4	4	В	0.932567	10723.1	4	В
0.957936	10439.1	1	В	0.932506	10723.8	1	В
0.957138	10447.8	4	В	0.932083	10728.7	5	В
0.956891	10450.5	4	В	0.931914	10730.6}	7 6	В
0.956616	10453.5	6	В	0.931611	10734.1 \$		В
0.955733	10463.2	5	В	0.930962	10741.6	7	В
0.955451	10466.3 }	4	B	0.9060	11038.		M(T)
0.955348	10467.4	6	В	0.9050	11050.		M
0.954393	10477.9	8	В	0.8230	12151.		M
0.953612	10486.4	5	В	0.8227	12155.		M(T)
0.952230	10501.7	10	В	0.7957	12568.		M(T)
0.951934	10505.0	4	B	0.77	12987		Many
0.951706	10507.4	ġ	B	0.7227	13837		M(T)
0.950171	10524.4)	8	B	0.7220	13850		M
0.950076	10525.5	ě	B	0.6994	14298		M(T)
0.949960	10526.8	4	B	0.6960	14368		M
0.949951	10520.6	6	B	0.69	14493		Many
0.949449	10532.4)	7	B	0.6530	15314		M
0.949341	10533.7	5	B	0.6524	15328		M(T)
0.948197	10546.3	9	B	0.6324	15813		M(T)
		-		0.5952	16801		
0.948023	10548.4	6	B	0.5932	16880		M(T)
0.946116	10569.5	9	B	0.5722	17476		M(T) $M(T)$
0.945996	10570.9	8	B B	0.3722	56040	1	Ra
0.945698	10574.2	1					<u> Ka</u>
0.945622	10575.0	5	В	0.17705	56480 }		Ra
0.945479	10576.6	4	B	0.17546	57000 {		
0.945412	10577.4	4	В	0.17392	57500 {		Ra
0.944339	10589.4	4	B	0.17153	58300 }		L
0.944089	10592.2	12	В	0.1700	58820		L
0.943790	10595.6	8	В	0.16948	58980 }		Ra
0.943068	10603.7	6	В	0.16805	59510 \		214
0.942836	10606.3 }	8	\mathbf{B}	0.16585	60300 (Ra
0.942685	10608.0	9	В	0.16421	60900 }		
0.9420	10616.		M(T)	0.16194	61750 }		Ra
0.941772	10618.3	6	В	0.15986	62560 \$		
0.941044	10626.5	6	В	0.15871	63010 }		D.
0.94	10638.		Many	0.15736	63560 {		Ra
0.938684	10653.2	9	В	0.15550	64310 }		Ra
0.938122	10659.6	9	B	0.15422	64850 \$		
0.937974	10661.3	4	В	0.1392	71840	1	L
0.937774	10663.5	9	B	0.13405	74600	1	Ra
0.937158	10670.6	10	В	0.13722	72880 }	W	Ra
0.936960	10672.8	7	B	0.13610	73470 {	• • •	214
0.936646	10676.4 (5	В	0.13567	73710)	w	Ra
0.936495	10678.1 յ	6	В	0.13462	74290 §	**	
0.935893	10685.0 }	7	В	0.13406	74590 }	1	Ra
0.935755	10 686.6 §	8	\mathbf{B}	0.13322	75060 (IXa
0.935526	10689.27	2	В	0.13242	75540 }		Ra
0.935450	10690.0 [6	В	0.13177	75890 §		1\a
0.935365	10691.0	4	В	0.13093	76380 (Ra
0.935306	10691.7	4	В	0.13063	76560 Š		лa
0.934567	10700.1	7	B	0.12973	77080 {		Ra
0.934405	10702.0 >	10	В	0.12910	77460 S		1/d
0.934252	ر 10703.8	7	В	0.12839	77890)		D.
0.933947	10707.2	6	В	0.12790	78180		Ra
0.933464	10712.8	6	В	0.12708	78690 (D.
0.933355	10714.0	7	B	0.12671	78920}		Ra
					_		

Table 62.—(Continued)

λ	$\boldsymbol{\nu}$	R^a	Ref.b	λ	v	R^a	Ref.b
0.12569	795 60 (Ra	0.10592	94410 (S	Ra
0.12531	79810 §		Na	0.10500	952405	J	ıta
0.12441	80380)		Ra	0.10430	95880 }	w	Ra
0.12428	80470	_1.		0.10409	960805		_
0.12407	80600	sh	Ra	0.10277	97310	Ssh	Ra
0.12381	80770	sh	Ra	0.10133	98690	S	Ra
0.12240 0.12141	81700 } 82370 {	S	Ra	0.10050 0.1000	99500 100000	s b	Ra H
0.12141	83800		Ra			b	11
0.11934	83940		Ra	0.09996 0.09984	100040 \ 100170 \	w	Ra
0.11977	834901			0.09960	1001703		Ra
0.11902	84020		Ra	0.09938	100400		
0.11769	849801		_	0.09917	100830		Ra
0.11701	85470		Ra	0.09811	101930	w	Ra
0.11520	86810)		_	0.0957	104500	ď	Ra
0.11486	87070		Ra	0.08568	116710	_	H
0.11334	88230		Ra	0.08506	117560		H
0.11293	88550)		Ra	0.08437	118520		H
0.11270	88730 \$	S	Na	0.08359	119630		H
0.11236	890001		Ra	0.08293	120580		H
0.11204	89250	w	Na	0.08218	121680		H
0.11164	89580)	s	Ra	0.08149	122710		H
0.11138	89790	5	144	0.07948	125820		H
0.11127	89870]	w	Ra	0.07841	127530		H
0.11118	89940 \$			0.07756	128930		H
0.10984	91040	sh	Ra	0.07667	130430		H
0.10951	91320	w	Ra	0.07588 0.07508	131790		H H
0.10911	91630	S	Ra		133190		
0.10883	91880	-		0.0745	134200	b	H Ra
0.10863 0.10839	92050 \ 92260 \	w	Ra	0.0740 0.0694	135100 144100	b b	Ra
0.10839	92800)		_	0.0590	169500	bs	Ra
0.10744	93080	S	Ra	0.0504	198400	bc	Ra
0.207 11	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			0.0001		20	

Remarks:

- region of continuous absorption begins here, and extends to shorter wave-lengths.
- beginning of the "continuum," which from here on underlies the lines and bands.
- strong absorption from here toward shorter wave-lengths, apparently bs made up of discrete bands (whether bs indicates anything essentially different from bc, is not clear).
- center of band.
- long wave-length limit of the band.
- very weak.
- weak. w
- strong.
- S very strong.
- sharp line.

Numerals indicate the intensity on a scale peculiar to the observer.

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- BC
- DH

Table 62.—(Continued)

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Witt, H., Z. Physik, 28, 236-248 (1924).

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K
L
M
Many
M(T)
McU
PIS
Ra
Ru<sup>1</sup>
  Ru<sup>2</sup>
RuA
    RuHt
  SPh
StW
WbR
Wi
WrR
```

From Paschen's observations v. Bahr inferred that water-vapor exhibits no selective absorption for $\lambda = 400 \,\mu$, approximately, and that there are bands at $\lambda = 250$ and 170 #.

Table 63.—Some Regions of Notable Transparency in the Absorption Spectrum of Water-vapor

E. K. Plyler and W. W. Sleator 357 have stated that there appears to be no absorption at the center $(3.168 \,\mu)$ of the band that is the harmonic of that near 6.26 µ.

Unit of $\lambda = 1 \mu = 10^4 A = 10^{-4} cm$

Region of transparency	Reference
115	H. Rubens. 358
91	358
75	358
62	358
47	358
3.168a	E. K. Plyler and W. W. Sleator.357
Between 1.38 and 1.87b	W. W. Sleator and E. R. Phelps. 359
Between 1.87 and 2.67b	359
Between 0.137 and 0.178c	G. Rathenau.337

- " See head of table.
- ^b Between these two bands there is no absorption comparable with that in the band at 1.38μ .
- Even at the highest pressure used there was a transparent region between the two bands beginning, respectively, at 0.137 \mu and 0.178 \mu.

and 65. M. Magat ³³⁰ has concluded that that analysis is certainly correct. R. Mecke ³³¹ has remarked: "It is very interesting that the normal frequency vibrating with a momentum parallel to the axis of symmetry does not occur in the absorption spectrum, although one would expect this one

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to have the strongest absorption. The reason why it does not absorb is not known."

The increase in the width of an absorption band when the pressure of the vapor itself is increased has been studied by S. D. Cornell,³³² who found for the half-width and its increase per atmosphere the values 0.57 \pm 0.06 cm⁻¹ and 0.13 cm⁻¹/atm for $\lambda = 0.94 \,\mu$, and 0.40 \pm 0.04 cm⁻¹ and 0.29 cm⁻¹/atm for $\lambda = 11.35 \,\mu$.

OH radicals are formed and excited when an electric discharge is passed through water-vapor; the excitation so produced gives the radical a relatively excessive amount of rotation, as is shown by its emission spectrum; its absorption spectrum shows only normal rotation.³³³ These excited radi-

Table 64.—Analyses of the Absorption Vibration Spectrum of Water-vapor

The most recent formulas for the computation of the normal frequencies of the water-vapor molecule are those derived by L. G. Bonner ³⁰⁰ from the positions found by R. Mecke ³⁶¹ for the band centers. He included terms in the fourth power of the distances and found $\omega_{\sigma} = 3796.0 - 39.5v_{\sigma} - 53.05v_{\pi} - 10.50v_{\delta}$, $\omega_{\pi} = 3674.8 - 53.05v_{\sigma} - 70.2v_{\pi} - 9.45v_{\delta}$, $\omega_{\delta} = 1615 - 10.50v_{\sigma} - 9.45v_{\pi} - 19.5v_{\delta}$.

The first close approximation to them was obtained by R. Mecke (Table 65); and a closer one by M. Magat, who obtained $\omega_{\sigma} = 3795 - 40v_{\sigma} - 50v_{\pi} - 10v_{\delta}$, $\omega_{\pi} = 3670 - 50v_{\sigma} - 71v_{\pi} - 10v_{\delta}$, $\omega_{\delta} = 1615 - 10v_{\sigma} - 10v_{\pi} - 19.5v_{\delta}$, which formulas he still considers the best. 330

Here the v's determine the harmonic; σ , π , δ indicate the types of vibration of the triangular molecule, σ is the "symmetrical" vibration, in which the O-atom oscillates along a line perpendicular to that joining the two H-atoms, π is the "antisymmetrical" vibration, in which the O-atom oscillates along a line almost parallel to that joining the two H-atoms, and δ is the "scissors" vibration, in which the II-atoms alternately approach and recede from one another. In every case, the other atoms oscillate in such

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0.63

0.60

0.59

0.57

Table 64.—(Continued)

a manner that the center of inertia of the molecule is unaffected by the oscillations.

The resulting wave number is given by the formula

$$v_0 = v_\sigma \omega_\sigma + v_\pi \omega_\pi + v_\delta \omega_\delta$$

The values of the v's assigned to each of the several band-centers, and the corresponding computed values for v_0 , are tabulated below.

	_			Bonner	Magat	Mecke	N	λ		
•	- υ· π	δ	Obs.		Computed		-	π	8	^
			600в				0	0	1	16.7
0	0	1	1595.5	1595.5°	1595.5	1595	0	1	0	6.30
0	0	2	3152.0	3152.0°	3151	3150	0	2	0	3.17
			35008				1	0	0	2.86
0	1	0	(3600)	3604.6						2.78
1	0	0	3756.5	3756.5°	3755	3756	0	2	1	2.66
1	0	1	5332.3	5331.0	5330.5	5331	1	0	3	1.88
1	1	0	7253.0	7255.0	7254	7247	1	2	1	1.38
1	1	1	8807.05	8810.6	8809.5	8802	2	0	3	1.14
1	2	0	10613.12	10613.1°	10612	10598				0.94
3	0	0	11032.36	11032.5°	11025	11034	•			0.91
1	2	1	12151.22	12149.8	12147	12133				0.82
3	0	1	12565.01	12565.0°	12560.5	12569				0.80
1	3	0	13830.92	13830.80	13826	13809				0.72
3	1	0	14318.77	14318.8¢	14324	14307				0.70
1	3	1	15347.91	15348.6	15340.5	15324				0.65

Unit of $\nu_0 = 1$ cm⁻¹; of $\lambda = 1$ $\mu = 10^4$ A = 10^{-4} cm.

15839

16817

16892

17493

15822

16799

16880

17445

3 1 1

1

1 4 0 15832.47

16821.61

16899.01

17495.48

15832.4°

16827.4

16908.1

17464.7

An alternative interpretation proposed by M. Magat ses to include the Raman bands; he states that it accounts for the observed spectrum as well as does Mecke's interpretation given in the first column.

^b Raman band.

[&]quot;Used in computing the constants.

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Table 65.—Molecular Constants Involved in the Vibration Spectrum of Water-vapor 329

 $v_0 = v_{\sigma}\omega_{\sigma} + v_{\pi}\omega_{\pi} + v_{\delta}\omega_{\delta}; \ \omega_{\sigma} = 3795 - 39(v_{\sigma} + v_{\pi}), \ \omega_{\pi} = 3670 - 70(v_{\sigma} + v_{\pi}), \ \omega_{\delta} = 1615 - 20(v_{\sigma} + v_{\pi} + v_{\delta}).$ The moments of inertia are I_{A} , I_{B} , I_{C} , related to the tabulated quantities, A, B, C, respectively, by means of the equations $I_{\Lambda} = h/8\pi^{2}cA = (27.658/A) \cdot 10^{-40} \text{ g·cm}^{2}$, etc., h being taken as 6.547×10^{-27} erg·sec, and c as $2.99796 \cdot 10^{10} \text{cm/sec}$; $10^{40}I_{A} = 0.996 + 0.045v_{\sigma} + 0.026v_{\pi} - 0.098v_{\delta}, \ 10^{40}I_{B} = 1.908 + 0.014v_{\sigma} + 0.033v_{\pi} - 0.034v_{\delta}, \ 10^{40}I_{C} = 2.981 + 0.047v_{\sigma} + 0.062(v_{\pi} + v_{\delta}) \text{ g·cm}^{2}; \Delta = I_{C} - (I_{A} + I_{B}); \ \alpha = \angle \text{ HOH}, \ r = \text{distance HH}, \ \text{H} \ \text{and O} \ \text{being the points occupied by the atoms H} \ \text{and O} \ \text{in the triangular water molecule}; \ v_{\sigma}, v_{\pi}, v_{\delta} \ \text{are quantum numbers}.$

Unit of $\lambda = 1\mu = 10^4 A = 10^{-4}$ cm; of $\nu_0 = 1$ wave/cm; of A, B, and C = 1 cm⁻¹; of $r = 1A = 10^{-8}$ cm; of $\Delta = 10^{-40}$ g·cm².

λ	2'σ 2'π 2'δ	m(obs.)	A	В	С	α	r	Δ
	000	0	27.81	14.50	9.284	105° 6′	0.970	0.077
6.296	001	1595.4	30.70	14.70	9.12	107° 30′	0.984	0.29
3.168	002	3152	35.8	15.0	9.0	111° 10′	0.984	0.46
2.663	100	3756.35	26.50	14.47	9.20	103° 45'	0.975	0.06
1.875	101	5332.3	29.0	14.6	8.8	106° 30′	0.995	0.30
1.379	110	7253	26.1	14.3	8.9	103° 40′	0.992	0.08
1.135	1 1 1	8807.0	28.60	14.7	8.71	105° 30′	1.002	0.33
0.9420	1 2 0	10613.25	25.25	13.89	8.75	103° 40′	0.998	0.075
0.9060	3 0 0	11032.33	24.45	14.18	8.87	102° 5′	0.991	0.032
0.8227	1 2 1	12151.23	27.84	14.17	8.66	105° 40′	1.001	0.248
0.7957	301	12565.01	26.75	14.41	8.74	104° 10'	0.997	0.212
0.7227	1 3 0	13830.91	24.70	13.71	8.61	103° 20′	1.005	0.075
0.6994	3 1 0	14318.73	24.04	13.95	8.69	102° 25′	1.004	0.050
0.6524	1 3 1	15347.90	27.15	13.97	8.52	105° 25′	1.012	0.247
0.6324	3 1 1	15832.47	(26.5)	(14.20)	(8.58)	104° 20′	1.013	(0.26)
0.5952	1 3 2	16821.62	29.28	14.03	8.34	107° 25′	1.023	0.400
0.5924	140	16898.81	25.02	13.60	8.50	103° 55′	1.014	0.115
0.5722	3 2 0	17495,44	23.72	13.84	8.32	102° 0′	1.026	0.160

20		0		1	1			2	
ขอ	υσ υπ υδ	α	100Δ	V	α	100Δ	V	α	100Δ
0 1 1 1	0 0 0 1 0 0 1 1 0 1 2 0 1 3 0	105° 6′ 103° 45′ 103° 40′ 103° 40′ 103° 20′	7.7 6 8 7.5 7.5	0 0 1 1 0 1 1 1 1 1 2 1 1 3 1	107° 30′ 106° 30′ 105° 30′ 105° 40′ 105° 25′	29 30 33 25 25	0 0 2	111° 10′ 107° 25′	46
1 3 3 3	1 4 0 3 0 0 3 1 0 3 2 0	103° 55′ 102° 5′ 102° 25′ 102° 0′	11.5 3.2 5.0 16.0	3 0 1 3 1 1	104° 10′ 104° 20′	21 26			

^a In their extended analysis (see p. 149), Randall and his associates used the following for the (0,0,0) values of A, B, and C: 27.8055, 14.499724, and 9.279276, respectively; these small changes from Mecke's values being made for the purpose of facilitating the computations.

Table 66.—Fine-structure of Absorption Bands of Water-vapor: Sources of Data

This table indicates the range in wave-length covered by the observations, the wave-lengths (λ) of the centers of the bands falling within that range, and where a report of the observations may be found. H. M. Randall ³³⁹ has reviewed the earlier data.

		Unit of $\lambda = 1\mu = 10000A = 10^{-4}$ cm.	
	Range:	Bands	Ref.
λ ₁	to λ ₂	λ	D
170	38		B W-D
135	60		WrR
75	18		R
24.9	10.13		WR
7.7	4.8	6.26	SP
7.6	4.8	6.26	$\mathbf{v}\mathbf{B}$
7.0	5.4	6.26	PIS, M
3.8	3.0	3.15	Ba
3.3	2.8	3.15	PIS, SP, M
2.8	2.5	2.67	PIS, S, M
2.65	1.35	1.37, 1.87, 2.67	Dr
1.9	1.8	1.87	PIS, SP, M, PI
1.42	1.35	1.37	SP, M, Pl
1.15	1.11	1.13	LH, M
0.96	0.93	0.942	BM, Br, LH, H
0.92	0.89	0.906	LH
0.92	0.78	0.906, 0.823, 0.796	\mathbf{BM}
0.73	0.71	0.723	\mathbf{BM}
0.71	0.69	0.699	FM
0.66	0.64	0.652, 0.632	\mathbf{FM}
0.63	0.56	0.595, 0.592, 0.572	FM
0.18	0.15		Ra, Pr
0.14	0.098		Ra, Pr

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Table 67.-Some Rotation Terms in the Infrared Spectrum of Water-vapor

(For other bands see references given in the text.) Mecke and associates. 329

11065.62 11099.92 11106.42 11122.45 11153.25 11163.46 11166.75 11198.26 11222.51 11229.65 11288.60 11244.55 11245.6 11296.70 11324.44 11331.58 11341.5 11342.8 11425.24 10687.33 10702.95 10737.05 10738.36 10743.60 10747.07 10777.43 10804.89 10811.45 10876.06 10823.87 10825.52 10883.15 10903.47 10911.18 10986.5 0921.67 0647.38 0680.43 11036.5 0652.44 0.6660 0.9420 8807.05 8830.45 8844.46 8850.36 8875.50 8885.21 8903.48 8944.85 8945.95 8940.13 8945.80 8979.6 9014.4 9020.7 (9103.5) 9022.49 9025.3 9079.8 9104.8 9121.9 9122.0 9123.2 9372.5 9239.1 λ_0 is the wave-length corresponding to $v_\sigma=v_\pi=v_\delta=0$ (see Table 65) 7253 7276 7290 7295 7321 7321 7815 1.379 7567 7683 7385 Unit of $\lambda = 1\mu = 10$ 4 = 10-4 cm; of $\nu = 1$ cm⁻¹ 5332.0 5355.4 5370.3 5376.6 5467.8 5473.8 5507. 5540.0 5547.8 5550.0 5553.5 5608.5 5635.5 5654 5401.7 5411.7 5428.6 5468. 5469. 5630. 5772 5653 5909 3896.1 3928.0 3956.9 3964.0 3975.5 3977.2 4025.7 4051.0 4064.2 4075.9 4030.0) 3756.5 3780.1 3792.1 3797.3 3834.1 3850.0 3886.1 3887.7 3892.0 3826.0 2.663 4195 4333 1732.3 1739.8 1772.7 1813.0 1820.2 1908.0 595.5 618.9 635.4 641.1 1665.3 1694.1 1742.5 1743.8 6.262 222.07 224.81 275.50 300.44 315.83 384.03 70.09 79.43 95.15 134.98 136.24 136.76 142.23 173.31 206.35 286.80 325.37 326.56 399.42 446.95 447.16 2

^{\$\\\ \}frac{1}{4} \cdot \frac{1}{4} \cdot \frac{1

		0.5722			17527.60																		17794.88	17002 00		•	
		0.5924	16899.01	16920.9	16932.20	16964.0	16971.3	16986.6	17023.1	17024.6	17029.1	17059.5	17088.4	17094.6	17167.2	17103.0	17105.3	17158.1	1/1/0.1	172604	17270.7		17199.01	17211 21	11011.6		
9523.7	9693.0 9878	0.5952	16821.61	16844.02	16859.26	16887.38	16898.25	16915.28	16960.64	16949.60	16956.22	16989.78	17026.53	17033.13	17110.72	17028.53	17033.27	17086.81	17114.41	1720073	17202.15		17128.50				
1969	8140 8328	0.6324	15832.47			15899.26		15922.21			15966.66																
6062	6238 6429	0.6524	15347.91	15370.50	15389.06	15413.94	15423.09	15439.45	15480.16	15476.22	15481.57	15514.01	15546.62	15552.93	15628.29	15556.87	15560.67	15610.99	15035.30	15051.79	15721.84		15652.43	15764 50	10101:03	15892.95	
4486	4663 4853	0.6994	14318.77	14341.39	14356.75	14385.03	14391.52	14407.25	14439.16	14447.20	14450.68	14481.70	14505.99	14512.91	14571.19	14526.37	14528.16	14578.10	14595.70	14612.93	14669.82		14622.07	14725 25	14133.33	14865.74	
		0.7227	13830.92	13853.25	13864.20	13896.46	13903.80	13919.05	13951.93	13958.17	13962.18	13992.61	14018.98	14025.58	14088.12 14088.50	14038.24	14038.26	14088.26	14110.1	14125.58	14202.2	14133.94	14134.2	0.50251			
		0.7957	12565.01	12588.16	12600.51	12632.88	12641.14	12658.20	12695.13	12696.74	12701.18	12734.88	12764.73	12771.76	12841.72	12777.51	12779.80	12834.43	12803.58	178/4.1/		12875.42	12876.70	12000 E	12991.0	13122.5	13272.1
744.1	920	0.8227	12151.22	12173.77	12187.75	12218.99	12227.82	12244.54	12286.79	12282.17	12287.20	12319.80	12353.80	12360.07	12437.11	12362.86	12365.22	12418.03	12445.81	12400.23	12531.68	12460.40	12461.32	17574 0	12574.0	12706.6	12855.4
e 0	9 - 8 0 - 10 9	<u></u>	0	1-1	- 1-	2-2	2-1	200	, z	3,1	3-2	3-1	30	ผู้เ	ئ م د	4	4-	4-8	4-1	of 4	F 4	5.4	7	7	7 7	7-1	9 9 0

cals last for at least 1/8 second after the discharge ceases.³³⁴ The absorption spectrum shows that these radicals are present also in the combustion zone of H_2 burning in O_2 (pressure 3 to 25 mm-Hg, furnace 470 to 550 °C), and in amount about 1000 times that corresponding to equilibrium.³³⁵

For a general discussion of the infrared spectra and for typical data obtained prior to 1930, see C. Schaefer and F. Matossi.³³⁶

In the far ultraviolet, water-vapor exhibits a continuous absorption from 1785 to 1550A; below 1550A there are several series of bands: one of doublets, $\Delta \nu = 170$ cm⁻¹, beginning at $\lambda = 1240$ A and accompanied by another displaced from it toward shorter wave-lengths by about $\Delta \nu = 3170$ cm⁻¹; another at 1220A accompanied by a weaker one displaced toward shorter λ 's by $\Delta \nu = 3240$ cm⁻¹; a third at 1091A; and a fourth at 1078A. These have been reported and analyzed by W. C. Price. The range 1700 to 957A had previously been studied by G. Rathenau, the with a smaller dispersion.

The ultraviolet portion of the spectrum is regarded as arising from changes in the electron configuration of the molecule; the near infrared bands, from the oscillations of the constituent atoms about their positions of equilibrium in the molecule; and the fine-structure of those bands, from rotations of the molecule as a whole, the effect of the rotation being superposed upon that arising from the oscillations of the atoms. In the far infrared are lines arising from the rotation alone. They are narrow when the pure vapor is at a low pressure, but broaden as the pressure is increased by the admission of air, owing to the impact of the molecules of air upon those of the vapor.³³⁸ As the H₂O molecule is triangular (see Section 9, Models) there are three principal axes of rotation, but only two of them seem to be spectroscopically effective.

An early report on the fine-structure of bands was published by H. M. Randall.³³⁰ The detailed analysis of the water-vapor spectrum rests upon the theoretical work of F. Lütgemeier ³⁴⁰ and of H. A. Kramers and G. P. Ittmann.³⁴¹ D. M. Dennison ³⁴² has reviewed the general subject; R. S. Mulliken ³⁴³ has studied and published the electronic configurations of the H₂O molecule; and in addition to those mentioned elsewhere, J. H. Van Vleck and P. C. Cross ³⁴⁴ have calculated from spectral data certain constants of the H₂O molecule. Certain relations connecting spectroscopic and thermal data have been considered by K. F. Bonhoeffer and F. Haber, ³⁴⁵ E. Justi, ³⁴⁶ and others. Their conclusions will be found in the sections devoted to the appropriate thermal data. In addition to those referred to elsewhere in this section, K. Basu, ³⁴⁷ G. Bosschieter and J. Errera, ³⁴⁸

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858 Rubens, H., Sitz. preuss. Akad. Wiss., 513-549 (1913).
850 Sleator, W. W., and Phelps, E. R., Astrophys. J., 62, 28-48 (1925).
850 Bonner, L. G., Phys. Rev. (2), 46, 458-464 (1934).
861 Mecke, R., Z. Physik, 81, 313-331 (1933).
862 Magat, M., Compt. rend., 197, 1216-1220 (1933).
863 Magat, M., Compt. rend., 196, 1981-1983 (1933).
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A. V. Buskovitch,³⁴⁹ and W. Weizel ³⁵⁰ have considered the interpretation of the spectrum of water-vapor.

The vibration of polyatomic molecules is the subject of a review by E. Bartholomé.⁸⁵¹

- 1. Band at $\lambda = 0.94 \,\mu$. The fine-structure of this band at 0.94 μ has been studied by F. S. Brackett (see Tables 62 and 66) and by J. H. Hsu.³⁵²
- 2. Region of $\lambda=18$ to 75 μ . The region 18 to 75 μ has been studied and analyzed with great care by H. M. Randall and his associates. They find that the rotational stretching and deformation of the molecule may correspond to a change of over 200 cm⁻¹ in the position of a line. They have determined the amount of this correction and have obtained good agreement between observed and computed λ 's up to quantum number j=12. For j=11 the vertex angle of the molecule is 98° 52′, and the O-H distance is 0.9640A, whereas the static values are 104° 36′ and 0.9558A. They have concluded that for components displaced much above 5 cm⁻¹ no first order correction for stretching, such as that proposed by E. B. Wilson, Jr., 354 can be satisfactory. They give extended tables of energy levels. A partial analysis of earlier data has been attempted by M. Eliaševič. 355
- 3. Region of 38 to 170 μ . The absorption of atmospheric water-vapor has been mapped from 38 to 170 μ and analyzed by R. B. Barnes, W. S. Benedict, and C. M. Lewis. The analysis was based on Mecke's values of the constants (Tables 64 and 65) and was carried, but very incompletely, to j=11. They concluded that the pure rotation spectrum of water-vapor is in complete agreement with the analysis.

Emission Spectrum.

The emission spectrum of water-vapor extends from the far infrared to the extreme ultraviolet, and consists of numerous bands, each containing many lines. Whether excited in the flame or by an electrical discharge, it is always accompanied by lines of hydrogen and of oxygen.

The ultraviolet bands have been studied in much detail. T. Heurlinger ³⁶⁴ found that most of the line of the 3064A band can be arranged in 12 branches forming what he called a band with doublet series; W. W. Watson ³⁶⁵ found that the 2811A band is of the same type, and concluded that both bands arise from the OH-molecule, formed in the flame and in the electric discharge. L. Grebe and O. Holtz ³⁶⁶ had previously suggested OH as the source of 3064. For details of the analyses of the bands, see E. C. Kemble, ³⁶⁷ D. Jack, ³⁶⁸ and especially R. S. Mulliken, ³⁶⁹ in addition to the authors cited in Table 68. R. T. Birge ³⁷⁰ has given certain con-

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<sup>301</sup> Huerlinger, T.. Diss., Lund, 1918.
<sup>805</sup> Watson, W. W., Astrophys. J., 60, 145-158 (1924).
<sup>806</sup> Grebe, L., and Holtz, O., Ann. d. Physik (4), 39, 1243-1250 (1912).
<sup>807</sup> Kemble, E. C., Phys. Rev. (2), 30, 387-399 (1927).
<sup>808</sup> Jack, D., Proc. Roy. Soc. (London) (A), 120, 222-234 (1928).
<sup>809</sup> Mulliken, R. S., Phys. Rev. (2), 32, 388-416 (1928).
<sup>870</sup> Birge, R. T., Int. Crit. Tables, 5, 415 (1929).
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Table 68.—Bands in the Emission Spectrum of Water-vapor

The emission spectrum of water-vapor extends beyond the limits covered by this table. F. Paschen ³⁷⁸ observed three bands of $\lambda > 8.2 \mu$, but was unable to determine their wave-lengths; and J. J. Hopfield ³⁷⁹ stated that the spectrum extends in the ultraviolet to about $900A = 0.09 \mu$. Each band contains many lines; each value of λ here given is that corresponding approximately to a maximum of emission, and serves to identify the band. In the infrared, the positions of the maxima are not exactly known, as the several determinations were made over 40 years ago, and differ among themselves.

 λ = wave-length, $\nu = 1/\lambda$ = wave-number, Bd = band designation in the paper first mentioned in the last column. With the exception of P and PWI, each of the cited papers includes a study of the fine-structure of the indicated band.

	Unit of $\lambda = 1\mu$	= 10^4 A = 10^{-4} cm; of ν	= 1 cm ⁻¹
λ	ν	Bd	Ref.a
8.2	1220		P
6.6	1510		P
5.6	1780		P P P
5.3	1880		P
2.8	3570		P
2.7	3700		N
1.8	5560		P, N
1.4	7140		P
0.3564^{b}	28060		RW1
0.3484	28700	1, 2	$_{ m JDW}$
0.3428	29170	0, 1	JDW, Ja
0.3328	30050	·	RW1
0.3185	31300	2, 2	DJ
0.3122	32030	1, 1	DJ, Ja
0.3064	32640	0, 0	DJ, Di, GH, Ja
0.3021	33100		$\mathbf{W}\mathbf{n}$
0.2875	34780	2, 1	DJ, Ja
0.2811	35570	1, 0	DJ, Wn, Ja
0.2676	37370	3, 1	CC
0.2608	38340		Ja CC
0.2447	40870	3, 0	CC

a References:

CC

 $[\]mathbf{DJ}$

Chamberlain K, and Cutter, H. B, Phys. Rev. (2), 44, 927 930 (1933). Dawson, D. H., and Johnston, H. L, Idem, 43, 980-991 (1933). Dieke, G. H., Proc. K. Akad. Wet. Amst., 28, 174-181 (1925); Nature, 115, 194 (1925). Di

GH

^{(1943).} Grebe, L., and Holtz, O., Ann. d. Physik (4), 39, 1243-1250 (1912). Jack, D., Proc. Roy. Soc. (London) (A), 115, 373-390 (1927); Idem, 118, 647-654 (1928). Ja Johnston, H. L., Dawson, D. H., and Walker, M. K., Phys. Rev. (2), 43, 473-480 JDW

Neunhoeffer, M., Ann. d. Physik (5), 2, 334-349 (1929); Idcm, 4, 352-356 (1930). Paschen, F., Idcm, (Wicd.), 50, 409-443 (1893); Idcm, 51, 1-39 (1894); Idem, 52, 209-237 (1894); Idem, 53, 334-336 (1894). Rodebush, W. H., and Wahl, M. H., J. Chem'l Phys., 1, 696-702 (1933); J. Am. Chem. Soc., 55, 1742 (L) (1933). Watson, W. W., Astrophys. J., 60, 145-158 (1924).

RWI

Degraded toward the red.

The band at 3064A seems to be associated with the formation of chemically "active gas" by the discharge.879a

stants of the OH-molecule, derived from spectral data. W. H. Rodebush and M. H. Wahl ³⁷¹ think that the bands at 3564A and 3328A arise from ionized OH.

The infrared emission bands have been studied mainly by Paschen. He found that the position of the maximum of a band shifts, and the intensity increases, as the temperature of the vapor is increased. The direction of the shift depends upon the band. He concluded that the emission probably obeys Kirchhoff's law. 872 He gives the following examples of the shift with temperature ³⁷³: $\lambda = 2.831$, 2.812, 2.717, and 2.661 μ , corresponding respectively to the temperatures 1460, 1000, 500, and 100 °C; $\lambda =$ 5.322, 5.377, 5.416, 5.607, 5.900, and 5.948 μ , corresponding respectively to the temperature of the oxy-hydrogen flame, 1470, 1000, 600, 100, and 17 °C; $\lambda = 6.620$, 6.597, 6.563, 6.527, and 6.512 μ , for 1470, 1000, 600, 100, and 17 °C. For the 2.8 μ and the 6.6 μ bands the maximum shifts to longer wave-lengths as the temperature increases; for the 5.3 \(\mu\) band the shift is in the opposite direction. E. v. Bahr 374 regarded the 6.6μ and 5.3 μ bands as components of a double band, the separation varying with the absolute temperature (T) in such a manner that $(\nu_2 - \nu_1)$ is proportional to \sqrt{T} ; and showed that Paschen's observations satisfied that relation. ν_1 and ν_2 being $1/\lambda_1$ and $1/\lambda_2$, respectively.

Over the range $6.254 \,\mu$ to $5.780 \,\mu$, and again at $5.188 \,\mu$, the emission is very weak, but it is strong at $5.636 \,\mu$, which lies between them.⁸⁷⁵

For a discussion of the occurrence of the OH radical in flames and in vapor through which a discharge has passed, see V. Kondratjew and M. Ziskin,³⁷⁶ T. Kitagawa,³⁷⁷ O. Oldenberg,³³³ and A. A. Frost and O. Oldenberg.³³⁴

A review of our knowledge of the vibrations of polyatomic molecules has been published by E. Bartholomé. 351

22. DIELECTRIC CONSTANT OF WATER-VAPOR

P. Debye ³⁸⁰ has shown that the dielectric constant of a gas composed of molecules that contain both permanent electrical dipoles and elastically bound electrons will satisfy relation (1)

$$\left(\frac{\epsilon - 1}{\epsilon + 2}\right) \frac{T}{\rho} = aT + b \tag{1}$$

³⁷¹ Rodebush, W. H., and Wahl, M. H., J. Am. Chem. Soc., 55, 1742 (L) (1933); J. Chem'l Phys., 1, 696-702 (1933).

⁸⁷² Paschen, F., Ann. d. Physik (Wied.), 51, 1-39, 40-46 (1894). See also Section 18.

⁸⁷⁸ Paschen, F., Idem, 51, 1-39 (1894); 53, 334-336 (1894).

⁸⁷⁴ v. Bahr, E., Verh. deut. physik. Ges., 15, 731-737 (1913).

⁸⁷⁵ Paschen, F., Ann. d. Physik (Wied.), 52, 209-237 (1894).

³⁷⁶ Kondratjew, V., and Ziskin, M., Acta Physicochim. URSS, 6, 307-319 (1937); 7, 65-74 (1937).

³⁷⁷ Kitagawa, T., Proc. Imp. Acad. Japan, 12, 281-284 (1936).

³⁷⁸ Paschen, F., Ann. d. Physik (Wied.), 50, 409-443 (1893).

³⁷⁹ Hopfield, J. J., Nature, 110, 732-733 (1922).

²⁷⁹a Urey, H. C., and Lavin, G. I., J. Am. Chem. Soc., 51, 3290-3293 (1929).

where ϵ = dielectric constant, ρ = density, T = absolute temperature, and a and b are positive constants characteristic of the gas. There are various reasons for believing that the molecules of water-vapor are of that kind. (See Section 9, Dipole moment.)

Table 69.—Dielectric Constant of Water-vapor (See also Figure 2)

For comments and references to original publications, see text. Debye's equation requires $(\epsilon - 1)T/\rho = 3(aT + b)$, approximately, a and b being positive constants characteristic of the gas or vapor. The best set of determinations is that of Stranathan, giving $3b = 3449 \pm 23$, $3a = 0.671 \pm 0.065$.

The investigator is indicated by his initial: $B = B\ddot{a}deker$, J = Jona, $SS = S\ddot{a}nger$ and Steiger, Z = Zahn.

Stranath	an. Comp	uted fro	m 3a = 0.	671, 3b =	= 3449.			
20 3.65	50 3.67	80 3.69	100 3.70	125 3.72	150 3.73	180 3.74	200 3.77	
10 ⁶ (€−1)			1		106(e-1)		-1) T 000ρ	
SS, $\rho = 0$ 400.2 371.7 348.8	3.76_{0} 3.76_{0}		23.3 29.6 39.6		25 23	(extrapolated) 3.8 ₆ 3.6 ₅ 3.2 ₄		
328.7 J, p =	3.794		107.9		15 12	3	3.7, 3.9,	
584 575	4.16 4.19		142.	2	765 767		5.90 5.98	
511 489	4.12 4.15 4.21		145.	8	736 694 648		5.76 5.50 5.21	
	$ \begin{array}{c} 20 \\ 3.65 \\ \hline $	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$					

As $(\epsilon + 2)$ is essentially equal to 3, and ρ is very nearly proportional to the pressure when T is invariable, relation (1) requires that $(\epsilon - 1)$ shall, for T constant, be essentially proportional to the pressure. Such proportionality for gases throughout the range 0 to 800 mm-Hg has been found by K. Wolf, ³⁸¹ using frequencies of 1 to 10 megacycles per second; but it could be tested for water-vapor only over a much shorter range in pressure, owing to difficulty in maintaining the insulation. But very recently, J. D. Stranathan ³⁸² has reported that he has found such proportionality to hold for water-vapor "to very near saturation," departures near saturation being due in many cases, perhaps in all, to the electrostatic polarization of the moisture adsorbed on the solid insulation, and not to leakage. From observations at 14 temperatures, ranging from 21.3 to 197.9 °C, he

⁸⁸⁰ Debye, P., Physik. Z., 13, 97-100, 295 (1912).

⁸⁸¹ Wolf, K., Ann. d. Physik (4), 83, 884-902 (1927).

found for the coefficients of equation (1) the values: $a = 0.2237 \pm 0.0217$, $b = 1149.6 \pm 7.8$. These are the best determinations now available:

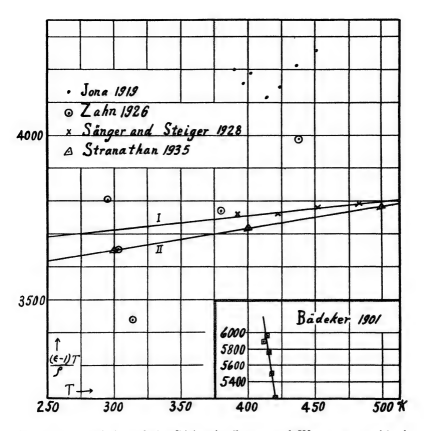


FIGURE 2. Variation of the Dielectric Constant of Water-vapor with the Temperature and Density.

Line 11, determined by Stranathan's equation (see Table 69), is probably the best available representation of the variation of $(\epsilon - 1)T/\rho$ with T. Line 1 closely represents the data obtained by Sanger and Steiger; its equation is $(\epsilon - 1)T/\rho = 3576 + 0.448T$. The three points (triangles) on line 11 were computed by means of Stranathan's equation; all other specially marked points represent experimental values. Badeker's data are totally at variance with the others. References are given in the text. $\epsilon =$ dielectric constant (cgse), ρ g/cm³ = density, T °K = absolute temperature.

The best previous data for the dielectric constant of water-vapor were those of R. Sänger and O. Steiger, 383 republished by Sänger, 384 and by R. Sänger, O. Steiger, and K. Gächter, 385

³⁸⁰ Stranathan, J. D., *Phys. Rev.* (2), **48**, 538-549 (1935) \rightarrow **47**, 794 (A) (1935), extending **45**, 741 (A) (1934).

³⁸⁸ Sänger, R., and Steiger, O., Helv. Phys. Acta, 1, 369-384 (1928).

⁸⁸⁴ Sänger, R., Physik. Z., 31, 306-315 (1930).

⁸⁸⁵ Sänger, R., Steiger, O., and Gächter, K., Helv. Phys. Acta, 5, 200-210 (1932).

Of the earlier determinations, those of C. T. Zahn, ³⁸⁶ were the best, and his results were used by H. L. Curtis in deriving the formula given in *International Critical Tables* ³⁸⁷ for the variation of ϵ with the temperature and the pressure. They are not inconsistent with those of Stranathan and of Sänger and Steiger, but are of such a low order of accuracy that the formula derived from them is quite erroneous. The abnormally rapid increase observed by Zahn in the apparent dielectric constant with the vapor-pressure when the pressure exceeded about a quarter of that for saturation, is probably not due to adsorption, as he supposed, but is more likely due to a failure in the insulation. ³⁸⁸ The determinations of M. Jona ³⁸⁹ are merely relative, the values of (ϵ – 1) being admittedly in error by an unknown factor. Those of K. Bädeker, ³⁹⁰ which are frequently quoted, seem to be seriously in error, as they impose a negative value upon the essentially positive coefficient a of equation (1), and are in other ways out of harmony with the results obtained by others.

All these sets of data are given in Table 69 and are displayed in Figure 2. The data available in 1926 have been discussed by O. Blüh,³⁹¹ who gives a bibliography of 172 entries.

23. DIELECTRIC STRENGTH OF WATER-VAPOR

Water-vapor is a perfect insulator* unless the strength of the applied electric field exceeds a certain critical value $(E_{\rm e})$ characteristic of the pressure (p) and the temperature. But near the region of saturation, condensation on solid insulators may destroy their insulating properties, and this phenomenon may be erroneously interpreted as an indication that the vapor itself is conducting in that region.

The quantity E_c has been called the electrical strength, the *elektrische Festigkeit*, and the *cohésion diélectrique*.³⁹² It differs from the sparking potential, in that the latter involves a term depending upon the electrodes, and is a function of the configuration of the spark-gap.

From observations in the range p = 0.05 to 5.4 mm-Hg, E. Bouty³⁰³ concluded that, at 22 °C, $E_c = 333 + 50.0p + 0.106/p^2$ volt/cm, the unit of p being 1 mm-Hg. This requires E_c to pass through a minimum (345 volt/cm) at p = 0.16 mm-Hg.

The over-all voltage (V_i) at which a current begins to pass between a given pair of electrodes, though obviously related to E_c , is not derivable

^{*}The conductivity arising from the action of radiations of various kinds is ignored in this section; it is relatively low in most cases.

⁸⁸⁶ Zahn, C. T., Phys. Rev. (2), 27, 329-340 (1926).

See Curtis, H. L., Int. Crit. Tables, 6, 78 (1929).
 Wolf, K., Physik. Z., 27, 588-591, 830 (1926); Ann. d. Physik (4), 83, 884-902 (1927). See however, Stranathan.

⁸⁸⁹ Jona, M., Physik. Z., 20, 14-21 (1919).

²⁰⁰ Bädeker, K., Z. physik. Chem., 36, 305-335 (1901).

⁸⁹¹ Blüh, O., Physik. Z., 27, 226-267 (1926).

²⁰² Bouty, E., Compt. rend., 131, 443-447, 469-471 (1900).

Bouty, E., Idem, 131, 503-505 (1900).

from it, and may differ from the sparking potential. It has been studied recently by E. Weichelt 304 and by S. Franck. 305 Using as electrodes brass spheres 4 cm in diameter, placed with their supporting rods and the gap along the axis of a cylindrical metal cage 12 cm in diameter, Weichelt found the ratio of V_4 for water-vapor to that for air, for the same gap and at the same temperature and pressure to be 1.16_6 . The gaps ranged from 0.5 to 3.2 cm, and temperatures from 115 to $160\,^{\circ}\text{C}$; the pressure was always 1 atm. The uncertainty of the individual measurements was about 2.5 per cent. For a given pressure, V_4 for air, and hence for water-vapor, varies inversely as the absolute temperature.

Table 70.—Voltage Difference at which Current Begins to Flow Through Water-vapor 395

See text for definitions and details.

Unit of $V_i = 1$ kilovolt; of gap = 1 cm

I. Comparison with air. Point and Plate.^a

Gap→ Gas	1.0	2.5	$5.0 - (V_{+}/V_{-})_{i}$	7.5	10
Vapor	1.100	1.057	1.057	1.014	1.015
Air	1.084	1.061	1.130	1.127	1.138

II. Vapor. Values read from graphs.

			- Point and Pla	ate		
Gap	1	2	4	6	8	10
$(V_+)_i$	4.6	4.9	5.2	5.4	5.6	5.7
$(V_{-})_{i}$	4.2	4.6	5.0	5.3	5.5	5.6
		Plates -			Spheres ^b	
V_{i}	0.66	1.33	2.0	0.5	1.0	1.5
V_{i}	18.8	33.9	48.7	14.1	27.3	37.4

[&]quot;The subscript + means that the potential of the point is positive with reference to that of the plate.

Franck, using a different type of apparatus and working at about 0.95 atm and about 100 °C, found V_i for water-vapor to be about 8 per cent greater than for air, the electrodes being either plates, or balls 5 cm in diameter; but if the gap is between a point and a plate, the ratio depends upon the length of the gap and upon whether the potential of the point is positive (V_+) or negative (V_-) with reference to that of the plate. If the gap exceeds about 3 cm, V_i for water-vapor is greater than that for air if the point is negative, and less if the point is positive. See Table 70.

The sparking potential (V_s) in water-vapor between clean spheres has been studied by I. Strohhäcker, ³⁹⁶ and that between a point and a plane by

^b Diameters = 5 cm.

⁸⁹⁴ Weichelt, E., Phys. Z., 32, 182-183 (1931).

²⁰⁵ Franck, S., Z. Phys., 69, 409-417 (1931).

⁸⁹⁰ Strohhäcker, I., Z. Physik, 27, 83-88 (1924).

S. Franck.³⁰⁵ The first used spheres 2 cm in diameter and worked at room temperatures and pressures not exceeding 15 mm-Hg. When his observed values of V_s are plotted against px, p mm-Hg being the pressure and x cm being the length of the spark-gap, a curve continuously concave to the axis of px is obtained for air. The same is true for water-vapor if px exceeds about 2.5, corresponding to $V_s = 700$ volts approximately; but between px = 2.5 and px = 1.5, V_s changes very little, and as px is reduced below 1.5, V_s decreases rapidly. The curve for the vapor cuts that for air near the point px = 5.8, $V_s = 890$.

Franck used a point made by sharpening a brass rod, 10 mm in diameter, to a cone with a vertex angle of 12°, and worked at 99 °C and a pressure of 728 mm-Hg. The opposing plate was earthed. The electrodes were illuminated with ultraviolet light. When the point is negative, $V_{\rm R-}=23x$ kilovolts for water-vapor, and 18.4x kilovolts for air, the potential for the vapor exceeding that for air by 25 per cent of the latter. When the point is positive the curve connecting $V_{\rm R+}$ and x is concave toward the axis of x, and for a given x, $V_{\rm 8+}$ for the vapor exceeds that for air by about 13 per cent of the latter. As read from the curve he gives, the values for water-vapor are as follows:

\mathcal{X}	1	2	3	4 cm
V_{s+}	14	25	34	40 kilovolts

An alternating voltage of 50 cycles/sec gave for the vapor almost exactly the same curve as was obtained with the point positive.

24. GLOW DISCHARGE IN WATER-VAPOR

(For the voltage and the field-strength at which conduction begins, see Section 23.)

Terms and Symbols.

Six distinct regions can be readily distinguished in the typical electrical discharge through a gas at low pressure and contained in a long tube with an electrode at each end. They will be designated as follows, beginning at the cathode:

- 1. Cathode glow. A thin layer of velvety glow closely adhering to the cathode.
- 2. Crookes dark space. A nonluminous region which increases in length as the pressure is decreased.
- 3. Negative column. A luminous region which is often called the negative glow. In this region and near the end that adjoins the Crookes dark space the electric field along the discharge becomes very weak, essentially zero as compared with the over-all voltage on the tube.
 - 4. Faraday dark space.
- 5. Positive column. A luminous region, in many cases striated, extending from the Faraday dark space, and by the shortest path, to the anode.

- 6. Anode glow. A thin glow closely adhering to the anode.
- (a) If the distance between the electrodes is progressively decreased, the pressure of the gas and the over-all voltage (V) on the tube remaining unchanged, the positive column suddenly vanishes when that distance has been reduced to a certain value (D), and at the same time V suddenly decreases by an amount ΔV which is generally equal to the ionization potential of the gas.
- (b) If d_o is the distance from the cathode to the boundary between the Crookes dark space and the negative column, d_o depends upon the pressure and V. If it exceeds a certain value $(d_o)'$ (about 0.7 mm in all cases), the application of a transverse magnetic field reduces it, but never below the value $(d_o)'$.
- (c) The distance (d_0) from the cathode to the point in the negative column at which the strength of the field becomes essentially zero is somewhat greater than d_c , though the two distances are often confounded.
- (d) The excess (V_{-}) of the potential at d_0 above that at the cathode is known as the cathode drop in potential. It does not differ greatly from the over-all voltage (V).

Table 71.—Vanishing of the Positive Column in Water-vapor

Adapted from A. Günther-Schulze 407 with the correction of certain arithmetical errors.

The increase in pD with p, when V=415 volts and p exceeds 1.47 mm-Hg, is attributed to the heating that occurs at those higher pressures.

D mm = Distance between the electrodes when the column just vanishes (\S a); p mm-Hg = pressure of the vapor; V volts = over-all difference in the potentials of the electrodes.

V = 415 volts			p = 1.30 mm-Hg				
Ď	D	pD	105D/V2	V	D	ÞĎ	$10^{6}D/V^{2}$
6.75	4.10	27.7	2.38	427	11.4	14.8	6.25
6.16	4.27	26.3	2.48	496	14.1	18.3	5.74
4.75	4.83	23.0	2.80	597	21.7	28.2	6.09
3.55	5.00	17.8	2.90	681	31.2	40.6	6.74
2.37	7.00	16.6	4.06	800	40.8	53.0	6.37
1.47	9.40	13.8	5.46			Mean	6.24
1.04	14.6	15.2	8.98				
0.828	15.2	12.6	8.83				
0.560	25.2	14.1	14.63				

Mean of last 4 = 13.9

Wherever in the remainder of this section it seems desirable to refer to one of the preceding paragraphs it will be done by writing in parentheses the number or letter designating the paragraph, preceded by the sign \(\xi \); thus (\(\xi \) c) will refer to paragraph (c) above.

³⁰⁷ Bonhoeffer, K. F., and Pearson, T. G., Z. physik. Chem. (B), 14, 1-8 (1931).

³⁸⁷a Brewer, A. K., and Kueck, P. D., Phys'l Chem., 38, 889-900, 1051-1059 (1934).

max Emeléus, K. G., and Lunt, R. W., Trans. Faraday Soc., 32, 1504-1512 (1936) (for Part II see Lunt).

Qualitative Relations.

In the glow discharge, water-vapor is dissociated and gives reactions characteristic of, but more active than, atomic hydrogen, reducing metallic salts not reduced by the latter.^{370a} The belief of Urey and Lavin ^{379a} that the OH molecule is present has been confirmed. It is present and excited, and

Table 72.—Effect of a Magnetic Field upon the Crookes Dark Space in Water-vapor 408

 d_c = distance from the cathode to the farther end of the Crookes dark space (see §§ b and c); p = pressure of the vapor; H = strength of the magnetic field transverse to the discharge.

Unit of $p=1$ mm-Hg; of $d_c=1$ mm; of $H=1$ gauss.							
$\stackrel{r\rightarrow}{H}$	1.00	3.00 de	19.23				
0	7.41	2.30	0.81				
120	3.77	-					
240	2.51						
360	1.55	1.54	0.61				
720	0.80	0.95	0.62				
1200	0.65	0.60	0.60				

Table 73.—Rate of Dissociation of Water-vapor in the Glow Discharge 405

I= current; v= volume of uncondensable gases produced per second (neither the pressure nor the temperature is stated); x= distance between the electrodes, which were plates 3 cm in diameter, the tube being 6 cm in diameter and 17 cm long. Observations were made at room temperature and a pressure of 0.75 mm-Hg.

v is not linear in I. Linder assumed that each H_2 in the gas corresponds to the dissociation of one H_2O . There was a slight deficiency of O_2 , only 30.0 to 30.4 vol. per cent being found.

Unit of $I = 1$ ma = 0.001 ampere; of $v = 1$ mm ³ /sec; of $x = 1$ cm.									
$\stackrel{x\rightarrow}{I}$	1.0	1.9	3.0	4.3	7.75	9.75			
1.0	0.239	0.159	0.166	0.150	0.173				
1.5			demonstrat	-	-	0.239			
2.0	-		•			0.343			
3.0	0.465	0.531	0.520	0.510	0.531	0.571			
5.0	0.717	0.822	0.916	0.862	0.929	0.909			
10.0	1.430	1.725	1.900	1.860	(2.020)1.925	2.230			
15.0	2.585	2.88	3.03	2.98		3.34			
17.0				-	3.61				
20.0	3.42	3.93₅	4.14	4.15	Name and A	\\\ 4.64 \\\ 4.36			
25.0	4.42	4.77	4.87	5.00	-	{5.97 {5.71			
26.0		******			5.53	(3,72			
28.0				5.62	-				

possesses an excessive amount of rotation.⁸³³ It can be detected spectroscopically for at least 1/8 second after the discharge stops ⁸³⁴; but the unexcited OH molecule has a very short life, of the order of 0.001 sec.⁸⁹⁷

The chemical reactions that occur in the glow discharge are numerous,

Table 74.—Distribution of Electrons in the Crookes Dark Space in Water-vapor 405

I= current, n= total number of electrons per second that cross a transverse section of the Crookes dark space at a distance x from the cathode, $n_0=$ total number of electrons per second that leave the cathode, $d_0=$ distance from cathode to place of zero field (see § c). Observations at room temperature and a pressure of 0.75 mm-Hg; electrodes, plates 3 cm in diameter in tube 6 cm in diameter; distance between electrodes = 3.5 cm.

Unit of $I=1$ ma = 0.001 ampere; of d_0 and $x=1$ mm.										
$\overset{I\rightarrow}{d_0\rightarrow}$	9.6	3 7.5	5 6.9	10 6.1 n/n	15 5.9 ₈	20 5.8 ₆	25 5.6			
1 2 3	1.96 3.77 6.82	2.07 4.36 8.42	2.26 4.39 9.35	2.22 4.86 10.50	2.43 5.42 11.80	2.25 5.21 11.30	2.34 5.50 12.60			
4 5 5.3	12.3 20.7	16.0 28.0	18.6 31.0	20.0 35.5	23.7 43.3	24.0 45.0	27.1 51.5			
6.0 6.5	32.8	45.0	49.0 56.0	52.0			57.0			
7.0 8.0 9.0	51.0 68.8 79.0	59.0								
d_{\bullet}	72.04	59.0	60.0	53.0	63.3	65.2	63.5			

[&]quot;This is obviously too small.

complex, and very sensitive to slight changes in the attendant conditions. A consideration of them lies beyond the scope of this compilation. Some of them are considered in the papers already mentioned; others in the following recent ones that have happened to come to the author's attention. In them references will be found to earlier work: A. K. Brewer and P. D. Kueck, ^{397a} K. G. Emeléus and R. W. Lunt, ³⁹⁸ K. H. Geib and P. Harteck, ³⁹⁹ K. H. Geib, ⁴⁰⁰ V. Kondratjew and M. Ziskin, ⁴⁰¹ R. W. Lunt, ⁴⁰² E. J. B. Willev. ⁴⁰³

Although the spectrum of the positive column in stationary water-vapor is the same throughout its length, II. O. Kneser 404 has observed that if the

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300 Geib, K. H., and Harteck, P., Idem, 30, 131-134, 140-141 (1934).
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⁴⁰⁰ Geib, K. H., J. Chem'l Phys., 4, 391 (L) (1936).

⁴⁰¹ Kondratjew, V., and Ziskin, M., Acta Physicochim. URSS, 5, 301-324 (1936).

⁴⁰² Lunt, R. W., Trans. Faraday Soc., 32, 1691-1700 (1936) (for Part I see Emeléus).

⁴⁰⁸ Willey, E. J. B., Idem, 30, 230-245-246 (1934).

⁴⁰⁴ Kneser, H. O., Ann. d. Physik (4), 79, 585-596 (1926).

⁴⁰⁸ Linder, E. G., Phys. Rev. (2), 38, 679-692 (1931).

⁴⁰⁰ Günther-Schulze, A., Z. Physik, 23, 334-336 (1924).

⁴⁰⁷ Günther-Schulze, A., Z. Physik, 30, 175-186 (1924).

vapor is streaming along the line of discharge, then, under suitable conditions, the spectrum of the positive column varies progressively from one end to the other, the intensity of the water bands, as compared with those of the Balmer lines and the continuous spectrum of hydrogen, decreasing

Table 75.—Dissociation of Water-vapor in the Glow Discharge See also Table 73. Adapted from E. G. Linder. 405

I= current, $d_0=$ distance from cathode to zero field (see § c), $V_-=$ cathode drop in potential (see § d), $n_1=$ number of electrons per second crossing the transverse section of the discharge at the distance d_0 from the cathode, $n_1W_1=$ total energy of those n_1 electrons, $n_1N=$ number of H_2 molecules formed per second, $k=W_1/N$. As k is found to be essentially constant, it follows that the number of molecules dissociated per second per electron entering the region of zero field is essentially proportional to the mean energy of those electrons.

Room temperature, p = 0.75 mm-Hg, electrodes were plates 3 cm in diameter and 3.5 cm apart in tube 6 cm in diameter.

Unit of $I=1$ ma = 0.001 ampere; of $d_0=1$ mm; of $V=1$ volt; of $W_1=1$ volt-electron/electron	on;
Unit of $I = 1$ ma = 0.001 ampere; of $d_0 = 1$ mm; of $V = 1$ volt; of $W_1 = 1$ volt-electron/electron of $N = 1H_2$ per electron; of $k = 1$ electron-volt per H_2 ; of $(V W_1)$ $V = 1$ per cent of V .	•

			-			$V W_1$
I	d_0	V	W_1	N	k	$\frac{V-V-V}{V-V}$
1	9.6	302	53.8	4.78	11.2	82.2
3	7.5	325	55.5	5.18	10.7	82.8
5	6.9	343	56.8	5.30	10.7	83.1
10	6.1	385	63.3	5.67	11.2	83.6
15	5.9₅	435	64.7	6.03	10.7	85.2
20	5.8₅	500	70.5	6.22	11.3	85.9
25	5.6	554	79.2	5.90	13.4	85.7
10 15 20	6.1 5.9 _δ 5.8 _δ	385 435 500	63.3 64.7 70.5	5.67 6.03 6.22	11.2 10.7 11.3	83. 85. 85.

Table 76.—Current-voltage Relation for Water-vapor between a Point and a Plane

Read from curves given by S. Franck. 395

Data for plate electrodes and low pressure are in Table 75.

Point was conical, vertex angle 12°, base 10 mm in diameter; plate was earthed; temperature 100 °C, pressure 725 mm-Hg.

I = current; x = distance from point to plane; V_* , $V_- = \text{potential of the point above, below, that of the plane.}$

Unit of I = 1 microampere; of x = 1 cm; of V = 1 kilovolt.

$x \rightarrow$		1		2.5		5.0	1	10
I	V_{+}	V	V_+	V_{-}	V_{+}	V-	V_{+}	V-
0ª	5.15	4.6	5.9	5.3	5.9			
10	6.9	6.8	10.2	10.6	14.5	13.6	17.9	17.0
20	8.2	8.8	13.1	14.0	19.2	18.5	23.2	24.0
30	9.4	10.3	15.6	16.4	22.1	22.1		
40	10.5	11.2		18.2		25.0		

^{*}These values for I=0 do not, but should, agree with those given in Table 70. The two sets of curves from which the values were read seem to be definitely discordant.

rapidly as the vapor advances along the tube. He regarded this as an indication that the emitters of these bands (probably OH ions) dissociate under the continued action of the discharge.

E. G. Linder 405 finds that almost all the dissociation of water-vapor occurs very near the cathode, probably within the distance d_0 (see § c). Hence, the actual amount of dissociation of OH ions in the positive column must be a very small part of the total dissociation of the vapor in the tube.

Numerical Data.

A. Günther-Schulze 406 has concluded that 85 per cent of the energy delivered to the cathode comes from the cathode drop in potential. This is confirmed by the more recent work of E. G. Linder. 405

Günther-Schulze 407 has concluded also that the thickness (d_+) of the anode glow is independent of the material used as anode and of the strength of the current. At such low pressures that d_+ is not greater than a small multiple of the mean free path of an electron, d_+ decreases as the pressure is increased; at higher pressures, d_+ is almost independent of the pressure. He gives the following values for water-vapor at room temperature:

Þ	1.11	2.66	3.55	14.45	mm-Hg
d_+	1.69	1.43	0.64	0.30	mm

The energy expended in producing the dissociation accompanying a glow discharge in water-vapor is 11 electron-volts per molecule of H_2O ionized (= 254 kcal/gfw = 1060 kj/gfw).⁴⁰⁵

IIB. WATER

25. MOLECULAR DATA FOR WATER

In this section are considered the structure of the liquid and the properties, or apparent properties, of the individual molecules, each regarded as identical with the molecule of the vapor.

Structure of Water.

The structure of the liquid will be considered under two heads: (1) Linkage, or association in its broadest sense, covering all types of bonding, temporary or permanent, between adjacent molecules of the vapor type; and (2) architecture, or the relative arrangement of the individual (vapor) molecules in a typical volume of the liquid or in a typical associated group, as the case may be.

Linkage of molecules.—On the basis of the kinetic theory of matter it is to be expected that collisions between dipole molecules will, in at least some cases, result in the colliding molecules remaining for a longer or shorter time close together and mutually oriented in a preferred manner, as a result of attraction between the dipoles; as has been suggested by

⁴⁰⁸ Günther-Schulze, A., Z. Physik, 24, 140-147 (1924).

J. Malsch ¹ and others. While so bonded they act as a single unit. These associated molecules may consist of two or more simple (vapor) molecules, and may be themselves bonded more or less strongly with other molecules, associated or simple, if all the molecules are crowded rather closely together, as in the case of a liquid. This bonding together of associated molecules may result in the formation throughout the liquid of numerous mutually independent groups of many molecules (hundreds or thousands), all in any one group having at any instant a similar orientation, but each group undergoing a relatively slow but continuous change in its personnel, size, and orientation. This has been called the cybotactic state.² Or, in the extreme case, each simple (vapor) molecule may be bonded to its nearest neighbors, and they to theirs, so that the entire volume of liquid forms, in a certain sense, a single loosely bonded molecule, the several individual bonds being continually broken and replaced by others. The distance between adjacent molecules will vary from point to point in the liquid. Where they are very close together the attraction between them will tend to keep them together with a preferred mutual orientation, thus welding them, at least temporarily, into an associated molecule; and such molecules may in turn be built temporarily into a definite architectural form characteristic of the substance. Thus the entire volume of the liquid may at any instant be quasicrystalline, the direction of the crystal axes varying from point to point, even over minute distances, and the entire picture changing from instant to instant.

All these views have been held. The last, the quasicrystalline concept, seems to be the most favored at present, especially by those engaged in the x-ray study of the structure of substances. It should be remembered that the duration of the structures so revealed need not be many times greater than the period of the x-rays used, provided that the structures are being continually renewed in form, but not necessarily either in the same place or with the same orientation. That period, of the order of 10⁻¹⁸ sec, is exceedingly short as compared with the mean time between molecular encounters in the liquid (even if the free path between encounters were as short as 0.001 of the mean distance between the centers of adjacent molecules, that time would be of the order of 10⁻¹⁵ sec). Consequently, the "structure" so revealed may be nothing more than an indication of the most frequent type of collision or of association, whether temporary or permanent.

Our knowledge and inferences regarding the molecular groupings and bondings to be found in water have been reviewed and discussed in a Symposium on the Constitution of Water 3 and by H. M. Caldwell, 4 G. G. Longinescu, O. Redlich, and T. C. Barnes and T. L. Jahn ; and more

¹ Malsch, J., Ann. d. Physik (5), 29, 48-60 (1937). ³ Stewart, G. W., Phys. Rev. (2), 35, 726-732 (1930).

⁸ Symposium, Trans. Faraday Soc., 6, 71-123 (1910).

⁴ Caldwell, H. M., Chem. Rev., 4, 375-398 (1927), bibliography of over 125 titles.

⁸ Longinescu, G. G., Idem, 6, 381-418 (1929).

⁶ Redlich, O., Monatsh. Chem., 53-54, 874-887 (1929).

briefly in a section of R. Kremann's article Wasser.8 The structure of liquids in general is the subject of a paper by J. Frenkel,9 of a symposium before the Faraday Society, 10 and of a paper by K. F. Herzfeld. 11 Individual views have been expressed by many, some of whom are mentioned in the following paragraphs. Both Caldwell and Kremann, especially the latter, have considered more particularly the evidence derived from various criteria for normality, such as, depression of the freezing point, elevation of the boiling point, ratio of the absolute temperature of the normal boiling point to that of the critical point, ratio of the molecular heat of vaporization to the absolute temperature of the normal boiling point (the increase in entropy on vaporization at the pressure of one normal atmosphere), the temperature coefficient of the molecular surface energy, etc.

Many types of observation (see Table 77) and in particular those showing that the behavior of water differs in very many respects from that of the large number of liquids commonly described as normal, suggest that water is not a simple substance, but is an equilibrium mixture of two or more interconvertible types, or groupings, of molecules, the composition of the mixture varying with the temperature and the pressure. variation is usually regarded as gradual, but E. J. M. Honigmann 12 is inclined to regard it as proceeding discontinuously.

The constituents of the mixture may conceivably be either isomeric or polymeric or both. Or they may be merely portions of the liquid in which the molecules are temporarily more closely packed or more uniformly oriented or both, than in adjacent portions. Or the abnormality may result from a uniform packing different from that in normal liquids. All these views have been advanced.¹³ At present, explanations in terms of packing and orientation seem to be the most favored,14 but the evidence is conflicting. For example, J. W. Ellis reports spectroscopic data which indicate a much closer union than any that can fairly be regarded as arising from a mere packing of the molecules; whereas Stewart draws the opposite conclusion from x-ray data.

Most of those who regard the abnormalities of water as arising from an association of molecules—from the presence of groups containing only a few molecules of H₂O each—seem to accept the opinion set forth in detail

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<sup>7</sup> Barnes, T. C., and Jahn, T. L., Quart. Rev. Biol., 9, 292-341 (1934).
      <sup>8</sup> Doelter, C., Handb. d. Mineralchem., 3, 855-915 (1918).
      <sup>9</sup> Frenkel, J., Acta Physicochim. URSS, 3, 633-648, 913-938 (1935).
     10 Symposium, Trans. Faraday Soc., 33, 1-282 (1937) → Nature, 139, 272-274 (rev.) (1937).
     11 Herzfeld, K. F., J. Appl. Phys., 8, 319-327 (1937), bibliography of 42 titles.
     <sup>12</sup> Honigmann, E. J. M., Naturwissenschaften, 20, 635-639 (1932).
<sup>18</sup> Longinescu, G. G., Chem'l Rev., 6, 381-418 (1929); Smyth, C. P., Idem, 6, 549-587 (1929); Stewart, G. W., Phys. Rev. (2), 43, 1426 (A) (1930); 37, 9-16 (1931); Indian J. Phys., 7, 603-615 (1932); Honigmann, E. J. M., Die Naturwiss., 20, 635-638 (1932); Bernal, J. D., and Fowler, R. H., J. Chem'l Phys., 1, 515-548 (1933); Fowler, R. H., and Bernal, J. D., Trans. Faraday Soc., 29, 1049-1056 (1933).
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½ Ellis, J. W., Phys. Rev. (2), 38, 693-698 (1931) → 38, 582 (A) (1931); Herzfeld, K. F.,
 J. Appl. Phys., 8, 319-327 (1937).
 ½ Röntgen, W. C., Ann. d. Physik (Wied.), 45, 91-97 (1892).

by W. C. Röntgen,15 but even then already old,* that water contains ice, that it is a saturated solution of ice in a liquid composed of simpler molecules. The most widely accepted assumption of this type seems to be that water is a mixture of (H₂O)₂ and (H₂O)₃, water-vapor not too near saturation being (H₂O), and the common type of ice (ice-I), not too near its melting point, being (H₂O)₈. In 1900, W. Sutherland ¹⁷ showed that many of the exceptional properties of water can be quantitatively accounted for on that assumption, and he derived the values of those properties that must be individually assigned to (H₂O)₂ and (H₂O)₃ in order to account for the observed values for water. These are given in Table 78. He proposed the names hydrol, dihydrol, and trihydrol, respectively, for the polymers (H_2O) , $(H_2O)_2$, and $(H_2O)_3$; these names are now quite generally used.

More recently, the idea that hydrol is present in water, especially at temperatures approaching the boiling point, has been advanced. This idea, that water contains hydrol (steam) in solution, is an essential part of H. L. Callendar's theory. 18 And it seems to be involved in the hypothesis by which C. Barus 19 sought to explain certain observations on the evaporation of very small drops near room temperature; viz., that water contains two constituents, one more volatile than the other.

But not a few who accept Röntgen's general suggestion take exception to Sutherland's interpretation; there is no general agreement as to the sizes of the individual groups of associated molecules. For example, G. B. B. M. Sutherland 20 accepts (H₂O)₂ but no higher polymer; C. Gillet 21 and H. E. Armstrong 22 have suggested that water is a mixture of (H2O) and (H₂O)₂; H. H. Vernon ²³ proposed (H₂O)₂ and (H₂O)₄; L. Schames ²⁴ and H. Auer 25 preferred (H₂O)₃ and (H₂O)₆, the first regarding ice as a mixture of $(H_2O)_6$ and $(H_2O)_{12}$. The more highly associated molecule of water was regarded as $(H_2O)_6$ by S. W. Pennycuick 26 ; as $(H_2O)_6$ or $(H_2O)_9$ by G. Tammann ²⁷; as $(H_2O)_6$, $(H_2O)_9$, $(H_2O)_{12}$, or $(H_2O)_{23}$ by J. Duclaux.²⁸ Some are noncommittal, e.g., C. S. Hudson,²⁹ A. Kling and A. Lassieur. 30

^{*} Röntgen's suggestion had been foreshadowed for at least 12 years. H. A. Rowland, in commenting upon the fact that the specific heat of water is a minimum at about 30 °C, had written: "However remarkable this fact may be, ... it is no more remarkable than the contraction of water to 4°. Indeed, in both cases the water hardly seems to have recovered from freezing. The specific heat of melting ice is infinite. Why is it necessary that the specific heat should instantly fall, and then recover as the temperature rises? Is it not more natural to suppose that it continues to fall even after the ice is melted, and then to rise again as the specific heat approaches infinity at the boiling point?" 16

¹⁶ Rowland, H. A., Proc. Am. Acad. Arts Sci., 15, 75-200 (131) (1880); "Physical Papers," 343-468 (398-399) (1902).

¹⁷ Sutherland, W., Phil. Mag. (5), 50, 460-489 (1900).

¹⁸ Callendar, H. L., Engineering (London), 126, 594-595, 625-627, 671-673 (1928); see also Jakob, M., Idem, 132, 651-653, 684-686, 707-709 (1931); Awbery, J. H., Rep. Prog. Phys. (Phys. Soc. London), 161-197 (1934).

Barus, C., Am. J. Sci. (4), 25, 409-412 (1908).
 Sutherland, G. B. B. M., Proc. Roy. Soc. (London) (A), 141, 535-549 (1933).

Neither is there agreement as to whether these more associated molecules exist as free individuals or form the blocks of a larger structural unit. Many regard them as free, but many others think that they are combined to form microcrystals of ice. H. Schade, 31 and H. Schade and H. Lofert 32 have advanced the idea that the polymers in water approach the size of colloid particles. But the light scattering units of which they are speaking may be merely regions where the molecules, as a result of their thermal agitation, are more closely packed than normal (see Section 39, Rayleigh scattering). H. T. Barnes 33 seems to think that as 0 °C is approached the (H₂O)₈ molecules in the water clump together to form invisible colloidal particles of ice; others think that they are temporarily associated into ill-defined groups of hundreds or thousands, all in any one group having essentially the same orientation,—the cybotactic state proposed by G. W. Stewart 34; and still others think that all the molecules are more or less closely bonded together into one quasicrystalline mass.

Of the quasicrystalline theories, three types have been proposed. One requires a fairly rigid structure; this was proposed by J. D. Bernal and R. H. Fowler 35 and in a modified form by M. L. Huggins at about the same time.³⁶ The other two types of theory are based on structures that are loose and very transient, 37 and will be considered in the paragraphs devoted to the architecture of water. Here it suffices to state that the Bernal-Fowler theory seems to be preferred at present. It is accepted. either directly or in a slightly modified form, by the workers here noted, 38-46 many of whom advance arguments in its favor.

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<sup>22</sup> Armstrong, H. E., Compt. rend., 176, 1892-1894 (1923).
<sup>23</sup> Vernon, H. H., Phil. Mag. (5), 31, 387-392 (1891).
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²⁴ Schames, L., Ann. d. Physik (4), 38, 830-848 (1912).

²⁵ Auer, H., Idem (5), 18, 593-612 (1933).

²⁰ Pennycuick, S. W., J. Phys'l Chem., 32, 1681-1696 (1928).

²⁷ Tammann, G., Z. physik. Chem., 84, 293-312 (1913); Z. anorg. allgem. Chem., 158, 1-16 (1926).

²⁸ Duclaux, J., Compt. rend., 152, 1387-1390 (1911); J. chim. phys., 10, 73-100 (1912).

²⁰ Hudson, C. S., Phys. Rev., 21, 16-26 (1905).

³⁰ Kling, A., and Lassieur, A., Compt. rend., 177, 109-111 (1923).

³¹ Schade, H., Z. Chem. Ind. Koll. (Koll. Z.), 7, 26-29 (1910).

⁸² Schade, H., and Lofert, H., Koll. Z., 51, 65-71 (1930).

⁸³ Barnes, H. T., in Alexander, J., "Colloid Chemistry," Vol. 1, pp. 435-443, Chemical Catalog Co., Inc. (Reinhold Publishing Corp.), 1926.

³⁴ Stewart, G. W., Phys. Rev. (2), 35, 1426 (A) (1930); 37, 9-16 (1931); Indian J. Phys., 7, 603-615 (1932).

³⁵ Bernal, J. D., and Fowler, R. H., J. Chem'l Phys., 1, 515-548 (1933); see also Fowler, R. H., and Bernal, J. D., Trans. Faraday Soc., 29, 1049-1056 (1933); Bernal, J. D., Idem, 33, 27-40-45 (1937): note E. Bauer's remarks (p. 43) about the last.

³⁰ Huggins, M. L., J. Phys'l Chem., 40, 723-731 (1936).

⁵⁷ Katzoff, S., J. (hcm'l Phys., 2, 841-851 (1934); Warren, B. E., J. Appl. Phys., 8, 645-654 (1937).

³⁸ Ananthakrishnan, R., Proc. Indian Acad. Sci., 2, 291-302 (1935).

³⁰ Cartwright, C. H., Nature, 135, 872 (L) (1935); 136, 181 (L) (1935); Phys. Rev. (2), 49, 470-471 → 421 (A) (1936).

⁴⁰ Cross, P. C., Burnham, J., and Leighton, P. A., J. Am. Chem. Soc., 59, 1134-1147 (1937).

⁴¹ Debye, P., Chem'l Rev., 19, 171-182 (1936).

V. Danilow ⁴⁷ departs from the Bernal-Fowler theory in some particulars; E. N. daC. Andrade ⁴⁸ is more inclined to favor the cybotactic theory advanced by Stewart; and P. Girard and P. Abadie ⁴⁹ think that the semi-

Table 77.—Evidence for Association

Some properties of water from which conclusions regarding an association of molecules have been drawn.

- 1. Magnetic susceptibility. The variation in the susceptibility with the temperature indicates the presence of at least two types of molecule, the more polymerized decreasing in number as the temperature rises.⁵¹⁻⁵⁷
- 2. Spectrum. The infrared spectrum of water indicates the presence of at least two types of molecule, their relative numbers varying with the temperature. At one time, J. W. Ellis and B. W. Sorge seemed inclined to accept the idea of polymerization, but later E. L. Kinsey and J. W. Ellis were undecided, rather inclining toward the quasicrystalline theory as propounded by Bernal and Fowler.
- 3. Raman effect: Scattering. It has been held that the Raman spectrum of water also indicates the presence of at least two types of molecule, their relative numbers varying with the temperature.⁶³⁻⁶⁹
- I. R. Rao concluded that there are three types of polymer involved (see Table 79). At one time, M. Magat ⁷⁰ seemed to favor the polymer theory, but he later decided ⁷¹ that such a view is untenable, and that a quasicrystal-

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42 Errera, J., J. de Chim. Phys., 34, 618-626 (1937).
   48 Herzfeld, K. F., J. Appl. Phys., 8, 319-327 (1937).
   44 Kinsey, E. L., and Ellis, J. W., Phys. Rev. (2), 49, 105 (L) \rightarrow 209 (A) (1936).
   45 Magat, M., Ann. de Phys. (11), 6, 108-193 (1936); Trans. Faraday Soc., 33, 114-120 (1937).
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<sup>51</sup> Piccard, A., Compt. rend., 155, 1497-1499 (1912); Arch. sci. phys. ct nat. (4), 35, 209-231, 340-359, 458-482 (1913).
   62 Cabrera, B., and Duperier, A., J. de Phys. (6), 6, 121-138 (1925).
   58 Mathur, R. N., Indian J. Phys., 6, 207-224 (1931).
   54 Johner, W., Helv. Phys. Acta, 4, 238-280 (1931).
   55 Auer, H., Ann. d. Physik (5), 18, 593-612 (1933).
   80 Azim, M. A., Bhatnagar, S. S., and Mathur, R. N., Phil. Mag. (7), 16, 580-593 (1933).
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   58 Collins, J. R., Phys. Rev. (2), 20, 486-498 (1922); 26, 771-779 (1925).
   50 Plyler, E. K., J. Opt. Soc. Amer., 9, 545-555 (1924).
   00 Ganz, E., Ann. d. Physik (5), 28, 445-457 (1937).
   en Ellis, J. W., and Sorge, B. W., J. Chem'l Phys., 2, 559-564 (1934).
   <sup>62</sup> Kinsey, E. L., and Ellis, J. W., Phys. Rev. (2), 49, 105 (L), 209 (A) (1936).
   68 Gerlach, W., Naturwissenschaften, 18, 68 (L) (1930).
   64 Meyer, E. H. L., Physik. Z., 31, 510-511 (1930).
   6 Specchia, O., Nuovo Cim. (N. S.), 7, 388-391 (1930).
   66 Segrè, E., Atti Accad. Linc. (6), 13, 929-931 (1931).
   67 Sutherland, G. B. B. M., Proc. Roy. Soc. (London) (A), 141, 535-549 (1933).
** Rao, I. R., Nature, 132, 480 (1933); Proc. Roy. Soc. (London) (A), 145, 489-508 (1934); Phil. Mag. (7), 17, 1113-1134 (1934).
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Table 77 .- (Continued)

line structure is required. R. S. Krishnan ⁷² could find in pure liquids no evidence of such large molecular clusters—comparable in size to a wavelength of light—as had been postulated by Plotnikow, ⁷³ but he did find such clusters in certain binary liquid mixtures.

- 4. Index of refraction.74
- 5. Color. J. Duclaux ⁷⁵ has suggested that the proportion of ice-molecules contained in water might be determined from a study of the thermal variation in the color. And H. T. Barnes ⁷⁶ has stated that "just at the freezing point the color of the St. Lawrence River changes rapidly and old river men can tell the approach of the ice-forming period by the color."
- 6. Diffraction of x-rays. The diffraction pattern obtained when x-rays are passed through water indicates that the water has a quasicrystalline character, changing with the temperature; but opinions differ as to whether the crystalline character refers to small groups of molecules or to large.⁷⁷⁻⁸⁴ For numerical data, see Tables 155 and 156.
 - 7. Dielectric constant.85, 86

86 Malsch, J., Physik. Z., 33, 383-390 (1932).

- 8. Surface tension.87
- 9. Pressure-volume-temperature correlations. G. Tammann ⁸⁸ has pointed out that water should contain at least as many types of molecule as there are types of ice that can exist in equilibrium with it. There were four such types then known that are stable (see Figure 13). Of these he concluded that III, V, and VI are isomers, all arising from molecules of the type (H₂O)₃. He gives reasons for believing that ice-I is either (H₂O)₉, breaking up into 9(H₂O), or is (H₂O)₆, breaking up into 2(H₂O)₃. From Bridgman's observations of the behavior of water under pressure, Tammann derived the values credited to him in Table 79. In the same table are the values derived by Jazyna and by Yoshida from the thermal expansion of water; they are much smaller than any of the others.

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    70 Magat, M., J. de Phys. (7), 5, 347-356 (1934).
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    <sup>72</sup> Krishnan, R. S., Proc. Indian Acad. Sci., 1, 211-216, 915-927 (1935); 2, 221-231 (1935).
    <sup>72</sup> Plotnikow, J., and Splait, L., Physik. Z., 31, 369-372 (1930); Plotnikow, J., and Nishigishi, S.,

Idem, 32, 434-444 (1931).
    74 Chéneveau, C., Compt. rend., 156, 1972-1974 (1913).
    18 Duclaux, J., Rev. gén. des Sci., 23, 881-887 (1912).

    <sup>76</sup> Barnes, H. T., "Ice Engineering," p. 10, Montreal, Renouf Publishing Co. (1928).
    <sup>77</sup> Keesom, W. H., and de Smedt, J., J. de Phys. (6), 4, 144-151 (1923); 5, 126-128 (1924).

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    81 Debye, P., and Menke, H., Physik. Z., 31, 797-798 (1930).
    82 Good, W., Helv. Phys. Acta, 3, 205-248, 436 (1930).
    88 Stewart, G. W., Phys. Rev. (2), 35, 726-732, 1426 (A) (1930); 37, 9-16 (1931).
<sup>84</sup> Kinsey, E. L., and Sponsler, O. L., Phys. Rev. (2), 40, 1035-1036 (A) (1932); Proc. Phys. Soc. (London), 45, 768-779 (1933).
    85 Errera, J., J. de Phys. (6), 5, 304-311 (1924).
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87 Ramsay, W., and Shields, J., J. Chem. Soc. (London), 63, 1089-1109 (1893).

Table 77 .- (Continued)

- E. E. Walker 89 and M. F. Carroll⁹⁰ have discussed the association of liquids in the light of van der Waals' equation of state and the concept of corresponding states, and D. B. Macleod 91 has considered certain phases of it from the concept of "free space."
- 10. Viscosity. From his measurements of the viscosity of water at two temperatures and under various pressures, P. W. Bridgman 92 concluded that there is an association that decreases as the pressure is increased, vanishing at very high pressures.
 - 11. Rate of cooling.93

Table 78.—Properties of Dihydrol and Trihydrol 17

$$2(H_2O) = (H_2O)_2 + 6.8 \text{ kcal} = (H_2O)_2 + 28.5 \text{ kilojoules.}$$

 $3(H_2O)_2 = 2(H_2O)_8 + 19.1 \text{ kcal} = 2(H_2O)_8 + 80.0 \text{ kilojoules.}^a$

	Te	mp.=0°C	
Property	$(H_2O)_2$	(H ₂ O) ₃	Unit
Density (p)	1.0894	0.88	1 g/cm^3
Coefficient of expansion $\left(-\frac{1}{\rho_0}\frac{d\rho}{dt}\right)$	9	2	10 ⁻⁴ (°C) ⁻¹
Specific refractivity $\left(\frac{n^2-1}{n^2+2},\frac{1}{\rho}\right)$	0.20434	0.20968	$1 \text{ cm}^3/\text{g}$
Compressibility $\begin{pmatrix} 1 & d\rho \\ \rho_0 & dP \end{pmatrix}$	16ª	10 (?)	$10^{-6} (atm)^{-1}$
Surface tension	78.3	73.3	1 dyne/cm °C
Critical temperature	368	538	°C.
Specific heat	0.84	0.6^a	1 cal/g·°C
Latent heat of fusion		16	1 cal/g
Latent heat of vaporization	257	250 ca.	1 cal/g
Viscosity (n)	3.0	38.1	10 ⁻³ poise
Pressure coefficient of viscosity		- 34	10 ⁻⁶ (atm) ⁻¹
Virial constant ^b	16.0	15.2	1 kiloatm cm ⁶ /g ²
Magnetic specific susceptibility	- 722.2	- 701.3	10 ⁻⁹ cgsm

^a J. Duclaux ^o concluded that the ice molecule is either $(H_2O)_0$ or $(H_2O)_{12}$, and was noncommittal regarding the formula for the "unpolymerized" water. He computed the following values, t=0 °C:

Heat of depolymerization of 1 g-molecule of dissolved ice = 4 kilocal. Specific heat of dissolved ice = $0.62 \text{ cal/g} \cdot ^{\circ}\text{C}$. Specific heat of "unpolymerized" water = $0.99 \text{ cal/g} \cdot ^{\circ}\text{C}$. Compressibility of "unpolymerized" water = $36 \times 10^{-6} \text{ (atm)}^{-1}$.

b By the virial constant he means the quantity l in the equation of state $\left(p + \frac{l}{2a^2}\right)v$ =RTf(v,T). Sutherland ⁹⁸ concluded that $f(v,T)=\frac{25}{13}\left\{1+\sqrt{T}\phi(v)\right\}$, whence $\frac{3}{4}l = v^2T\left(\frac{dp}{dT}\right)_a + \frac{25}{26}vRT - \frac{3}{2}pv^2$, the last term being negligible. He gives five methods for estimating the value of l for a liquid.

^e Derived by L. Sibaiya [∞] for water at 20 °C; for H₂O he finds -775.5 in the same units.

Table 79.—Estimates of the Ice-content of Water

If the average molecule of the mixture can be represented by $(\dot{H}_2O)_{\beta}$, β may be called the average degree of association of the mixture. O. Fruwirth 97 has tabulated 10 ways for getting β , the values ranging from 1.3 to 4.7. It is probable that β is the quantity called "degree of association" by M. A. Azim, S. S. Bhatnagar, and R. N. Mathur 98 for which they give the values 4.18 at 0 °C and 2.70 at 100 °C. It is not stated how the values were obtained. By the method of Ramsay and Shields (see p. 520), J. Timmermans and H. Bodson 99 derive $\beta=3.0$ from their own observations, probably near room temperature.

c = total mass of all the ice molecules contained in a unit mass of water at $t \, {}^{\circ}\text{C}$; n = number of molecules of the indicated type per 100 molecules of the mixture.

Unit of $p = 1 \text{ kg*/cm}^2 = 0.968 \text{ atm}$; of P = 1 atm.

I. Pressure = 1 atmosphere.

Ref ⁿ →	Su	Du	Ta 10	0c	Rao	Y06	·	Miscellaneo	Ref.
0	37.5	18.3	14.6	39.0	37	0.6	0	55	Sch
4					32	0.5	0	39	RC
10		14.3	10.4			0.3	0	1.70	Ja
20	32.1	11.3	7.0	26.2		0.2	0	34	Jo
30		9.1	4.4			0.1	0	24	Pi
38					32		10	1.00	Ja
40	28.4	7.4	2.5	15.4		0.05	20	28	ŘC
60	25.5	5.1		7.8		0.01	20	0.20	Ja
80	23.4	3.7		3.4		0	20	30	Au
98					21		20	15	Go
100	21.7	2.8		0			20	28	Jo
							30	Oc	Ja
$\underset{t}{\text{Type}} \rightarrow$	H ₂ O	10	(H ₂ O) ₂ 00¢ (Rao)	(H ₂ C))3	H ₂ O	(H n (l	(20)2 Rao)	(H ₂ O) ₃
0	9		57	3	7	19	5	8	23
4	10		58	3:		19.5		8.5	22
38	16		52	3:		29	5		21
98	21		58	2		36	5		13
Ice, 0 °C	0		32	68	8	0	4	1	59

II. Variation with pressure.

<i>f</i> →	1	500	1000	1500 c(Ta)	2000	2500
- 10	20.2	14.5	13.1	9.6	6.4	5.3
0	14.6	10.0	7.8	5.6	4.2	3.9
10	10.4	7.1	5.7	4.1	3.1	3.0
20	7.0	4.7	3.9	2.8	2.1	2.1
30	4.4	2.8	2.4	1.7	1.4	
40	2.5	1.4	1.3	0.7	0.7	0.8

 ⁶⁸ Tammann, G., Z. physik. Chem., 84, 293-312 (1913); Z. anorg. Chem., 158, 1-16 (1926).
 Walker, E. E., Phil. Mag. (6), 47, 111-126, 513-525 (1924).

Table 79.—(Continued)

$\stackrel{t\rightarrow}{P}$	0	20	40	0¢(Su) 60	80	100
1 150 $-10^6 \Delta c/\Delta P$	37.5 35.1 160	32.1 30.0 147	28.4 26.4 133	25.5 23.7 120	23.4 21.7 107	21.7 20.3 93
Extreme est	imates at	0 °C (D.				

Extreme estimates at 0 °C (Du).

\boldsymbol{P}	0	100	500	1000	1500	2000	2500
100 Cmax	18.3	16.9	12.2	8.1	5.4	3.6	1.6
100 Cmin	18.3	16.8	11.8	7.3	4.0	2.2	0.5

* References:

- Au Go
- Auer, H., Ann. d. Physik (5), 18, 593-612 (1933).
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- RC
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 Yoshida, U., Mem. Coll. Sci. Kyoto, 19, 271-277 (1936). Re Sch Su
- Ta Yo
- ^b Computed by means of his formula: $c = 0.006e^{-0.0044t}$ based on a study of the thermal variation of the density and of the specific heat.

crystalline state is well developed in water at room temperature, but do not commit themselves to the Bernal-Fowler theory. On the other hand, S. A. Ukholin 50 thinks that certain Bernal-Fowler interpretations of the thermal variations in the Raman spectrum of water are of quite doubtful validity.

Establishment of equilibrium.—If water is a mixture of two or more types of molecular structure, the relative proportions of the several types varying with the temperature, then the question arises: Does an appreciable time elapse after a change in temperature or in phase before the types are again in mutual equilibrium?

H. T. Barnes 100 has advanced the idea that equilibrium takes place slowly, at least at temperatures near 0 ° C, and has described experiments

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90 Carroll, M. F., Idem (7), 2, 385-402 (1926).
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[&]quot;These values were derived from the thermal expansion of water.

⁹¹ Macleod, D. B., Trans. Faraday Soc., 21, 145-150, 151-159 (1925).

⁹² Bridgman, P. W., Proc. Amer. Acad. Arts Sci., 61, 57-99 (1926).

⁹⁸ Vernon, H. M., Phil. Mag. (5), 31, 387-392 (1891).

⁹⁴ Duclaux, J., J. de chim. phys., 10, 73-109 (1912).

⁸⁵ Sutherland, W., Phil. Mag. (5), 35, 211-295 (241) (1893).

⁹⁶ Sibaiya, L., Current Sci., 3, 421-422 (1935).

or Fruwirth, O., Sitz.-ber. Akad. Wiss. Wien, Abt. (IIb), 146, 157-167 (1937) = Monatsh. f. Chem., 70, 157-167 (1937).

⁹⁸ Azim, M. A., Bhatnagar, S. S., and Mathur, R. N., Phil. Mag. (7), 16, 580-593 (1933).

⁹⁹ Timmermans, J., and Bodson, H., Compt. rend., 204, 1804-1807 (1937).

¹⁰⁰ Barnes, H. T., Scientific Monthly, 29, 289-297 (1929).

¹⁰¹ Barnes, T. C., and Jahn, T. L., Proc. Nat. Acad. Sci., 19, 638-640 (1933); Quart. Rev. Biol., 9, 292-341 (1934).

interpreted as showing that water may be temporarily deprived of the nuclei -assumed to be the ice-molecules in it—required for the formation of ice; and T. C. Barnes and T. L. Jahn 101 have reported experiments interpreted as showing that water freshly formed from steam freezes less quickly than that freshly formed by melting ice, the initial temperature and other conditions being the same for each specimen of water. The conclusion is drawn that an appreciable time is required for equilibrium to be established. In view of the uncertainty of our knowledge of the behavior of water as regards freezing, all such interpretations must be accepted with reservations. T. C. Barnes and associates 101, 102 have described experiments indicating that the growth of certain organisms in water freshly formed from steam differs from that in water freshly formed by melting ice. 102 The explanation offered attributes this difference to a difference in the icecontent of the two kinds of water; that is, these workers believe that a very appreciable time is required for the establishment of equilibrium.* Inspired by this work, A. P. Wills and G. F. Boeker 103 interpreted in the same way certain peculiar observations they obtained in their study of the magnetic susceptibility of water, but they found later that the peculiar behavior was the result of errors. 104 J. Zeleny 105 invoked such delay as an explanation of certain observations on the electrification of water-drops broken by an air-blast.

Other attempts to answer the question have led to the conclusion that equilibrium is attained very quickly, practically at once, some of the properties investigated being these:

- (1) Vapor-pressure. 106-110
- (2) Magnetic susceptibility. 111-113
- (3) Infrared absorption. 114
- (4) Index of refraction. 115
- (5) Density. Using a method by which a difference of one in 106 in the density could be detected, M. Dole and B. Z. Wiener ¹¹⁶ compared the
- *Such a conclusion is quite at variance with certain observations made by the compiler after this section had been written (see Section 97).

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<sup>102</sup> Barnes, H. T. and T. C., Nature, 129, 691 (1932); Barnes, T. C., Proc. Nat. Acad. Sci., 18, 136-137 (1932); Lloyd, F. E., and Barnes, T. C., Idem, 18, 422-427 (1932); Barnes, T. C., and Larson, E. J., J. Am. Chem. Soc., 55, 5059 (1933).
     103 Wills, A. P., and Boeker, G. F., Phys. Rev. (2), 42, 687-696 (1932).
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104 Wills, A. P., and Boeker, G. F., Idem, 46, 907-909 (1934).
105 Zeleny, J., Idem, 44, 837-842 (1933).
100 Bonhoeffer, K. F., and Harteck, P., Z. physik. Chem. (B), 5, 293-296 (1929).
107 West, W. A., and Menzies, A. W. C., J. Phys'l Chem., 33, 1893-1896 (1929).

106 Wright, S. L., and Menzies, A. W. C., J. Am. Chem. Soc., 52, 4699-4708 (1930).
100 Menzies, A. W. C., Proc. Nat. Acad. Sci., 18, 567-568 (1932).

110 Egerton, A., and Callendar, G. S., Phil. Trans. (A), 231, 147-205 (A698) (1932).
111 Auer, H., Ann. d. Physik (5), 18, 593-612 (1933).
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¹¹² Cabrera, B., and Fahlenbrach, H., Z. Physik, 82, 759-764 (1933).

LaMer, V. K., and Miller, M. L., Phys. Rev. (2), 46, 907-909 (1934).

LaMer, V. K., and Miller, M. L., Phys. Rev. (2), 43, 207-208 (1933).

¹¹⁶ Dole, M., and Wiener, B. Z., Science (N. S.), 81, 45 (1935).

density of freshly condensed steam with that of freshly melted ice, and found no difference.

In most cases, the conditions were not identical with those in the experiments of T. C. Barnes and his associates, water heated for a time to 100 °C being used instead of condensed steam, and water that had remained long at a much lower temperature, instead of freshly melted ice. Although this difference might be expected to reduce the magnitude of the effect sought, it would scarcely make the effect entirely negligible as compared with that corresponding to the other conditions.

G. Tammann and A. Elbrächter¹¹⁷ attempted to answer the question by comparing the observed temperature change accompanying adiabatic expansion with that computed from the specific heat and the equation of state. Differences were observed, but were of such a kind that they could not be interpreted.

Other information regarding the controversy may be found in articles by T. C. Barnes, ¹¹⁸ W. D. Bancroft and L. P. Gould, ¹¹⁹ and by A. W. C. Menzies, ¹²⁰ See also pp. 638, 644.

At one time H. B. Baker ¹²¹ thought that he had evidence that equilibrium between the molecular species in a pure liquid might be established slowly. However, it seems that his conclusions are not accepted. ¹²²

F. Vlès ¹²³ has explained the appearance of a Tyndall cone of light scattered by "cold water from freshly melted very pure ice" as due to colloidal aggregates of trihydrol. Before this explanation can be accepted it must be shown that it did not arise from foreign particles suspended in the liquid, and dissolving slowly as the temperature rose.

Certain observations briefly reported by F. W. Gray and J. F. Cruickshank ¹²⁴ as indicating that the thermal change in the magnetic susceptibility of water lags behind the temperature should be carefully studied and checked by other observers.

The Earl of Berkeley ¹²⁵ has suggested that information regarding association might be obtainable from a study of changes produced in the index of refraction as the liquid is centrifuged.

Architecture of water.—It has long been known that oxygen behaves as if it had 4 valencies, two weaker than the others, and hydrogen as if it had two, one being weak (see, e.g., T. M. Lowry and H. Burgess). Making use of the first, H. H. Vernon 23 suggested that, between 4 and 100 °C, water consists of two H₂O units joined by a double bond between the O's; and below 4 °C of 4 H₂O units arranged in a square and united

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<sup>117</sup> Tammann, G., and Elbrächter, A., Z. anorg. allgem. Chem., 200, 153-167 (1931).
<sup>118</sup> Barnes, T. C., Science (N. S.), 79, 455-457 (1934); 81, 200-201 (1935).
<sup>119</sup> Bancroft, W. D., and Gould, L. P., J. Phys'l Chem., 38, 197-211 (1934).
<sup>120</sup> Menzies, A. W. C., Science (N. S.), 80, 72-73 (1934).
<sup>121</sup> Baker, H. B., J. Chem. Soc. (London), 121, 568-574 (1922); 1927, 949-958 (1927).
<sup>128</sup> Menzies, A. W. C., J. Phys'l Chem., 33, 1893-1896 (1929); Science (N. S.), 80, 72-73 (1934).
<sup>129</sup> Vlès, F., Chem. Abst., 31, 592 (1937) ← Arch. phys. biol., 13, 199-201 (1936).
<sup>124</sup> Gray, F. W., and Cruickshank, J. F., Nature, 135, 268-269 (L) (1935).
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125 Earl of Berkeley, Nature, 120, 840-841 (1927).

by single bonds between adjacent O's. H. E. Armstrong 127 regarded water as a mixture of H₂O ("hydrone") and of (H₂O)₂ formed by uniting an H and an OH, each individually, to the O of an H₂O. This type of (H₂O)₂ he called "hydronole." A. Kling and A. Lassieur ³⁰ pictured water as a mixture of H₂O and a series of its polymers, regarding H₂O as existing in two isomeric forms: H—O—H, and H—OH. The suggestions of Armstrong and of Kling and Lassieur have been criticized by V. Auger. 128

If the 4 pairs of electrons of the oxygen atom are arranged about the nucleus at the vertices of a regular tetrahedron, 129 then the simple molecule H₂O will be formed by adding an H to each of two vertices of that tetrahedron; the other two vertices may for convenience be called the electron vertices of the H₂O molecule. A pair of such tetrahedral molecules may unite, as a result of the attraction between the H vertices of one and the electron vertices of the other, to form a double molecule in any one of three ways: (a) by union at a single vertex (type V); (b) by union at the two vertices of an electron-H edge (type E); and (c) by union at the three vertices of a single face (type F), the one electron and two H vertices of a face of one uniting, respectively, with the one H and two electron vertices of a face of the other. Such tetrahedral molecules can be built up into structures that correspond to the observed x-ray pattern of ice, and serve as the basis of much of the recent discussion regarding the structure of water and of ice.

- F. O. Anderegg, ¹³⁰ and E. L. Kinsey and O. L. Sponsler ¹³¹ suggested a face union (F). The former regarded such double molecules as forming the chief molecular species in water, and the latter regarded them as the polymer that occurs in both ice and water, dissociation being into H+ and $(H_3O_2)^{-}$.
- S. W. Pennycuick¹³² discussed the structure of water, especially in relation to such tetrahedral molecules, and concluded that the single vertex union (V) is the most probable. This not only yields a satisfactory crystal form, but also gives rise to rings (H₂O)₆ and to open chains. See also M. L. Huggins. 138

The x-ray pattern for both ice and water indicates that, on the average, each oxygen atom has four or very nearly four others as near neighbors. Taking account of this in connection with the tetrahedral molecule. I. D.

¹²⁰ Lowry, T. M., and Burgess, H., J. Chem. Soc. (London), 123, 2111-2124 (1923).

¹²⁷ Armstrong, H. E., Compt. rend., 176, 1892-1894 (1923).

¹²⁰ Smyth, C. P., Phil. Mag. (6), 47, 530-544 (1924); Pennycuick, S. W., J. Phys'l Chem., 32, 1681-1696 (1928).

¹⁸⁰ Anderegg, F. O., Chem. Abstr., 18, 2282 (1924) ← Proc. Indiana Acad. Sci., 1923, 93-101 (1923).

¹⁸¹ Kinsey, E. L., and Sponsler, O. L., *Proc. Phys. Soc.* (London), 45, 768-779 (1933) \rightarrow *Phys. Rev.* (2), 40, 1035-1036 (A) (1932).

¹⁸² Pennycuick, S. W., J. Phys'l Chem., 32, 1681-1696 (1928).

¹²⁸ Huggins, M. L., J. Phys'l Chem., 40, 723-731 (1936) → J. Am. Chem. Soc., 58, 694 (L). (1936).

Bernal and R. H. Fowler ¹³⁴ developed their theory of the quasicrystalline structure of water, which seems to be the favored one today. The exact crystal structures of ice and water are not determined unambiguously by the available data: there is an element of choice. They decided that three types of structures, passing continuously one into another, have to be considered: Type I is like tridymite; Type II is quartz-like; Type III is close-packed, like ammonia. Type I is ice and occurs in water at low temperatures; Type II predominates in water between 0 and 100 °C; Type III characterizes water at high temperatures, say between 150 °C and the critical point. There is no question of different kinds of molecules, but only of different arrangements of the same kind. Each small region of water has instantaneously a crystalline character, but in different regions the crystals are differently oriented, and each region is continuously changing in personnel and in crystal orientation. See also J. D. Bernal. ¹³⁵

- B. E. Warren ¹³⁶ does not admit as close or permanent a binding of the atoms into crystal forms as is postulated by Bernal and Fowler. He regards the crystal form as merely a kind of ideal that is more or less closely approached in water at any instant, but that is seldom, if ever, fully realized.
- S. Katzoff ¹³⁷ goes still further. He thinks that in water there is little if any periodicity in arrangement, and that what little there may be is entirely incidental. In his view, the important thing is the relative positions of adjacent molecules. They are probably held together in nearly the same manner as in the crystal, but except for that, the arrangement of the molecules is a random one. He found no evidence either for the definite "quartz-like" arrangement or for the extensive degree of close packing postulated by Bernal and Fowler; his observations were, in fact, incompatible with the assumption of a "quartz-like" arrangement. His proposed picture is that of a broken-down ice structure.

The earliest models placed the bonding H midway between the two O's, but it is probable that it is nearer to one than to the other, and that, of the 4 H's bonding to an O, two are nearer than the other two. $^{133, 134}$ And there are reasons for thinking that in H_2O one H is bound more firmly than the other (see L. Henry, 138 and G. Jacoby 139).

A. Piekara ¹⁴⁰ has pointed out that the rotating of a dipole molecule is affected by fields arising from two sources: the field due to its immediate neighbors and called by him the association field; and the resulting field arising from all the other, more distant, molecules, which he calls the Debye molecular field.

¹⁸⁴ Bernal, J. D., and Fowler, R. H., J. Chem'l Phys., 1, 515-548 (1933); Fowler, R. H., and Bernal, J. D., Trans. Faraday Soc., 29, 1049-1056 (1933).
¹⁸⁵ Bernal, J. D., Trans. Faraday Soc., 33, 27-40-45 (1937); and E. Bauer's remarks (p. 43).
¹⁸⁶ Warren R. E. J. Appl. Phys. 8, 645-654 (1937).

¹⁸⁰ Warren, B. E., J. Appl. Phys., 8, 645-654 (1937).
187 Katzoff, S., J. Chem'l Phys., 2, 841-851 (1934).

¹⁸⁸ Henry, L., Bull. Classe Sci. Acad. Roy. Belg., 1905, 377-393 (1905).

¹⁸⁸ Jacoby, G., Ann. d. Physik (4), 72, 153-160 (1923).

Piekara, A., Acta Phys. Polon., 6, 130-143 (1937).
 Callendar, H. L., Proc. Roy. Soc. London (A), 120, 460-472 (1928); Proc. Inst. Mechan. Eng., 1929, 507-527 (1929).

Miscellanea.—Both H. L. Callendar ¹⁴¹ and O. Maass and A. 1.. Geddes ¹⁴² find that the liquid structure may persist at temperatures appreciably above the critical one—above that at which the meniscus vanishes. (Cf. Section 88, Critical data.)

Evidence thought to show that the structure of water in capillary spaces differs from that of water in bulk has been published by P. Gaubert, who found birefringence in the film between two adjacent bubbles, and by E. Torporescu, who observed certain voltaic effects. The observations reported by B. Derjaguin so indicating that thin films of water possess a rigidity that increases as the film becomes thinner, conflict with the observations of R. Bulkley said and are probably to be otherwise explained, as it is not certain that Derjaguin had satisfactorily eliminated the effect of small suspended particles.

A. P. Wills and G. F. Boeker ¹⁴⁷ infer from their magnetic measurements that "significant changes in molecular arrangement or association of the water molecules" may occur near 35 °C and near 55 °C. And M. Magat ¹⁴⁸ concludes that many properties of water have anomalies near 40 °C, and regards these anomalies as arising from a change in the structure of water.

Dipole Moment of the Molecule of Water.

The value of the dipole moment (μ) of a single free molecule of H₂O as it occurs in water-vapor is such that $10^{19} \mu = 18.3_1$ cgse, see p. 48. A discussion of Debye's dipole theory, his formulas, and their limitations in the case of liquids may be found in Section 49.

All observations combine to show that Debye's formulas for freely reorientable dipoles, though applying to gases, are not applicable to dipole liquids, and are only approximately applicable to dilute solutions of dipole substances in nonpolar solvents. Nevertheless, those formulas have been quite generally used for determining what has been called the dipole moment of liquids. The procedure followed has been this: To the experimentally determined values of $(\epsilon - 1)/(\epsilon + 2)\rho$ is fitted an expression of the general form a + b/T + cT, sometimes with c = 0, and from this the apparent dipole moment denoted by μ_n , is determined by means of the relation $b = 4\pi N(\mu_a)^2/9Mk$. In these equations ϵ is the dielectric constant, ρ the density, T °K the absolute temperature, N the number of molecules per gram-mole (6.061×10^{23}) , k the molecular gas constant $(1.372 \times 10^{-16} \text{ erg/°K·molecule})$, and M the formula weight, which is assumed to be the molecular weight of the molecule to which μ_a

¹⁶² Maass, O., and Geddes, A. L., Phil. Trans. (A), 236, 303-332 (1937).

¹⁴⁸ Gaubert, P., Compt. rend., 200, 304-306, 679-680 (1935).

¹⁴⁴ Torporescu, É., *Idem*, **202**, 1672-1674 (1936) = Bull. de Math. et Phys., Bucarest, **6**, 40-41 (1936).

¹⁴⁵ Derjaguin, B., Z. Physik, 84, 657-670 (1933); Phys. Z. Sowj., 4, 431-432 (1933).

¹⁴⁰ Bulkley, R., Bur. Stand. J. Res., 6, 89-112 (RP264) (1931).

¹⁴⁷ Wills, A. P., and Boeker, G. F., Phys. Rev. (2), 46, 907-909 (1937).

¹⁴⁸ Magat, M., J. de Phys. (7), 6, 179-181 (1935); Trans. Faraday Soc., 33, 114-120 (1937).

refers. This quantity (μ_a) is what has been generally called the dipole moment of the liquid, and has been denoted by μ . Here it will be called the apparent dipole moment, and will be denoted by μ_a , μ being used to denote the dipole moment of a molecule of the gas phase. The ratio $(\mu_a/\mu)^2$ is the factor by which the theoretical value of b for freely reorientable molecules must be multiplied in order to obtain the value appropriate to the liquid. Its value is commonly indicated by that of μ/μ_a , the square root of its reciprocal.

The value to be preferred at present for the coefficient b for water is 107.13 °K·cm³/g (see Table 175),

```
10^{19} \mu_a = 5.59 cgse units per gfw-H<sub>2</sub>O
whence
                  \mu_a = 0.117_1 electron angstroms per gfw-H<sub>2</sub>O
which gives \mu/\mu_a = 3.27.
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This and other values that have been published for μ_a are given in Table 80. The value 1.7 given by L. Kockel was obtained by fitting to her

Table 80.—Apparent Dipole Moment of Water.

The apparent dipole moment (μ_q) is here used to denote the value derived from the coefficient (b) of T^{-1} in the expression for $(\epsilon - 1)$ $(\epsilon + 2)\rho$ in powers of T, by means of the relation $b = 4\pi N(\mu_q)^2/9kM$, M being the formula weight of H₂O (see text). Introducing the numerical values of the several quantities, one finds for water 1.850(10¹⁹) $\mu_a = \sqrt{b}$. For water-vapor c = 0 and μ_a in the expression for b is the actual dipole moment (μ) of the molecule H₂O (10¹⁹ μ = 18.3₁ cgse, p. 00).

Unit of $\mu_a = 10^{-19}$ cgse un	its $(=0.02$	1094 electron angstrom) per gfw-H ₂ O
μа	μ/μ_a	Reference (see text for comments.)
5.59	3.27	Preferred values.
5.7	3.21	P. Debye 151
5.689	3.218	M. Forró 152
1.7	10.8	L. Kockel 153
7.4	2.47	P. Lertes 154

own observations Debye's expression for gases (that containing only the a and b terms), although her observations demand the c term also; and P. Lertes' value, 7.4, was derived from the torque exerted upon water by a rotating electric field. Although both these values are included in the table it is probable that little, if any, weight should be attached to them.

Values of μ_a as derived from solutions of water in nonpolar solvents may be found, with references, in the table compiled largely by N. V. Sidg-

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149 Hampson, G. C., and Marsden, R. J. B., Trans. Faraday Soc., 30, Appendix (1934).
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¹⁵⁰ Frank, F. C., Proc. Roy. Soc. (London) (A), 152, 171-196 (1935).

¹⁸¹ Debye, P., Physik. Z., 13, 97-100, 295 (1912).

¹⁸⁹ Forró, M., Z. Phys., 47, 430-445 (1928). 188 Kockel, L., Ann. d. Physik (4), 77, 417-448 (1925). 184 Lertes, P., Z. Phys., 4, 315-336 (1921); 6, 56-68 (1921).

¹⁸⁶ Cennamo, F., Nuovo cim., (N. S.), 13, 304-309 (1936).

wick and published by G. C. Hampson and R. J. B. Marsden, ¹⁴⁹ and such results have been discussed by F. C. Frank ¹⁵⁰ and others.

Polarizability of the Molecule of Water.

By the polarizability of a molecule of a substance is meant that portion of the molecular electric moment that is induced per unit electrical field as a result of the relative displacement of elastically bound electrons. It is the quantity α occurring in Debye's formula $(\epsilon - 1)/(\epsilon + 2)\rho = (4\pi N/3M) \cdot (\alpha + \mu^2/3kT) = a + b/T$, [see Section 49, eq. (2)], and is related to the optical index of refraction n in accordance with the formula $a = 4\pi N\alpha/3M = (n^2 - 1)/(n^2 + 2)\rho$. For water (M = 18.0154), this gives $10^{24} \alpha = 7.096a = 7.096(n^2 - 1)/(n^2 + 2)\rho$.

The value for a that is to be preferred at present is 0.2262 cm³/g (see Table 175), which gives $10^{24} \alpha = 1.605$ cgsc, or $\alpha = 1.605$ (angstrom)³. On the other hand, using the values of n for the D-line, as given in Table 135 and of ρ as given in Table 93, one finds for 10, 20, and 30 °C the following values: $\alpha = 1.462_8$, 1.462_1 , and 1.461_7A^3 , respectively. F. Cennamo 155 obtained, from his own values of n, values essentially agreeing with these; it should be noticed that the values he gives are of α , not of α . The dielectric constant of water vapor leads to $\alpha = 1.59A^3$ for the vapor molecule (p. 49); and the optical index, to 1.50 (p. 49).

Anisotropy of the Molecule of Water.

For a discussion of the anisotropy of molecules, an explanation of terms and symbols, and references to the general subject, see Section 9, Anisotropy, and J. W. Beams. 156

If the electrical moments induced along the principal electrical axes of the molecule by a unit electrical field parallel to those axes are A, B, and C, respectively, and if the magnetic moments similarly induced along the same axes are A', B', and C', then the factor (δ) measuring the optical anisotropy

of the molecule is
$$\delta = \frac{A^2 + B^2 + C^2 - (AB + BC + CA)}{(A + B + C)^2}$$
 and the mean

susceptibility per molecule of the unmagnetized substance is $\theta' \equiv (A' + B' + C')/3$. It is generally assumed that B = C, and that B' = C', A being the greatest of the three induced electrical moments. Then $\delta = \left(\frac{A-C}{A+2C}\right)^2$, $\theta' = (A' + 2C')/3$, and C'/A' serves to define the magnetic

anisotropy; C'/A' is commonly called the magnetic anisotropy, although its value is unity for an isotropic molecule and zero for extreme anisotropy.

The data given by M. Ramanadham 157 lead to the following values, that of δ being taken from I. R. Rao 158 and based on the observations of

¹⁸⁸ Beams, J. W., Rev. Mod. Physics, 4, 133-172 (1932).

¹⁸⁷ Ramanadham, M., Indian J. Phys., 4, 15-38 (1929).

¹⁸⁸ Rao, I. R., Idem, 2, 61-96 (1928).

¹⁸⁰ Krishnan, K. S., Phil. Mag. (6), 50, 697-715 (1925).

¹⁶⁰ Chinchalkar, S. W., Indian J. Phys., 6, 165-179 (1931).

K. S. Krishnan ¹⁵⁹: $\delta = 0.00553$, $A'/\theta' = 1.14$, C'/A' = 0.81 if B' = C'. These values rest upon his value (-1.1×10^{-14}) for the coefficient of magnetic birefringence, and that is numerically greater than the more recent values $(-0.3_8 \times 10^{-14})$. From the latter, S. W. Chinchalkar ¹⁶⁰ computes $A'/\theta' = 1.05$, whence C'/A = 0.93 if B' = C'.

The value of δ for water is only 1/3 as great as that (0.0166) for water-vapor (cf. p. 50).

26. Interaction of Water and Corpuscular Radiation

Alpha Particles.

The range of alpha particles in water (liquid) at 15 °C is given by W. Michl ¹⁶¹ as 32 microns for rays from Po, and by H. R. v. Traubenberg and K. Philipp ¹⁶² and K. Philipp ¹⁶³ as 60 microns for rays from RaC'. (See S. Meyer. ¹⁶⁴)

The decomposition of water by alpha particles, and the nature of the products formed, have been studied by C. E. Nurnberger. 165

Beta Rays.

The coefficient of mass absorption of water (liquid) for the β rays from Ra-E is given by G. Fournier ¹⁶⁶ as $\mu/\rho = 17.4$ cm²/g; that calculated by him from the absorption by H₂ and O₂ on the assumption that the coefficients of atomic absorption are not affected by the union of the atoms to form molecules is 16.0. He regards the difference as an evidence that water is abnormal.

For luminescence excited by β rays, see Section 39, Electron Luminescence.

Neutrons.

The coefficient of absorption of neutrons in water is $\mu = 0.027$ cm⁻¹.¹⁶⁷ A table based on Fermi's expression for the slowing down of neutrons by water has been published by G. Horvay.¹⁶⁸

The scattering of neutrons by water has been studied by J. R. Dunning and G. B. Pegram ¹⁶⁹ and by M. Deisenroth-Myssowsky, I. Kurtschatow, G. Latyschew, and L. Myssowsky. ¹⁷⁰

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161 Michl, W., Sitz. Akad. Wiss. Wien (Abt. IIa), 123, 1955-1963, 1965-1999 (1914).
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¹⁶² v. Traubenberg, H. R., and Philipp, K., Z. Physik, 5, 404-409 (1921).

¹⁶⁸ Philipp, K., Idem, 17, 23-41 (1923).

¹⁰⁴ Meyer, S., Int. Crit. Tables, 1, 367-369 (1926).

¹⁶⁸ Nurnberger, C. E., J. Phys'l Chem., 38, 47-69 (1934); J. Chem'l Phys., 4, 697-702 (1936).

¹⁰⁶ Fournier, G., Compt. rend., 183, 200-203 (1926).

¹⁶⁷ Arzimowitsch, L., Kurtschatow, I., Latyschew, G., and Chramow, W., *Physik. Z. Sowj.*, **8**, 472-486 (1935). In this article the name of the last author is incorrectly spelled "Chromow"; its corrected later. They do not state the unit used for μ , but it is probably the one (cm⁻¹) here given.

¹⁶⁸ Horvay, G., Phys. Rev. (2), 50, 897-898 (1937).

¹⁶⁹ Dunning, J. R., and Pegram, G. B., Phys. Rev. (2), 45, 768-769 (A) (1934).

¹⁷⁰ Deisenroth-Myssowsky, M., Kurtschatow, I., Latyschew, G., and Myssowsky, L., Physik. Z. Sowj., 7, 656-669 (1935).

TENSILE RUPTURE OF WATER

Although it has long been known that liquids can, under suitable conditions, sustain relatively great tensile loads, it is doubtful if a true tensile rupture of a liquid is observable, unless it is when a liquid is forced to flow at high speed through a constricted section of a tube, as described by O. Reynolds 171 before the British Association for the Advancement of Science, in 1894. In all ordinary cases, the breaking of a column of liquid proceeds by a process of constriction arising from the action of surface tension; and when a column of liquid gives way under the action of a direct tension, the failure appears to occur at the liquid-solid boundary, not in the liquid itself unless that is known to contain a dissolved gas, in which case the failure seems to be associated with the presence at that point of a bubble of gas. Unless the principal radii of curvature (r_1, r_2) of the bounding surface are so small that the concept of an invariable surface-tension is not validly applicable to it, no element of liquid abutting upon a gas can sustain a tension exceeding $T\left(\frac{1}{r_1} + \frac{1}{r_2}\right)$, T being the surface-tension. For exam-

ple, if it is valid to apply the idea of an invariable surface-tension to bubbles 0.00001 cm (= 1000Å) in diameter, then the presence of a bubble of air of that size when under the tension will, of itself, cause a column of water to break at a tension of about 30 atm, however much greater the true tensile strength of water might be. On the other hand, the effect of completely dissolved gas upon the tension to which water can be practically subjected seems to be negligible, as shown particularly by the observations of M. Berthelot, ¹⁷² H. H. Dixon and J. Joly, ¹⁷³ and H. H. Dixon. ¹⁷⁴ Gas may also be the main cause of the observed failures at the liquid-solid boundary. All the recorded work was done before the advent of modern methods for outgassing solid surfaces. (See F. Donny, 175 O. Reynolds, 176 and J. Meyer. 177)

Two procedures, mainly, have been used for determining the tension to which water may be experimentally subjected. One is based upon the well known sticking of the mercury column to the top of a barometer when the tube and mercury are clean; it involves the determination of the length of the mercury column that can be similarly supported from a thin layer of water adhering to the top of the tube. This has been used by O. Rey-

¹⁷¹ Reynolds, O., "Papers on Mechanical and Physical Subjects," Vol. 2, pp. 578-587, Cambridge Univ. Press, 1901.

¹⁷² Berthelot, M., Ann. de chim. ct phys. (3), 30, 232-237 (1850).

¹⁷⁸ Dixon, H. H., and Joly, J., Phil. Trans. (B), 186, 563-576 (1895).

¹⁷⁴ Dixon, H. H., Proc. Roy. Dublin Soc., 12, 60-65 (1909); Ann. Report Smithsonian Inst. for 1910, 407-425 (1911).

¹⁷⁶ Donny, F., Ann. dc chim. ct phys. (3), 16, 167-190 (1846) = Ann. d. Physik (Pogg.), 67, 562-584 (1846).

¹⁷⁶ Reynolds, O., "Papers on Mechanical and Physical Subjects," Vol. 1, pp. 394-398, Cambridge Univ. Press, 1900.

Meyer, J., Abh. deuts. Bunsen-Ges., 3, No. 1, whole No. 6, 1911 = "Zur Kenntniss des negativen Druckes in Flüssigkeiten," W. Knapp, Halle, 1911.
 Reynolds, O., Ibid., pp. 230-243, 394-8.

¹⁷⁹ Moser, J., Ann. d. Physik (Pogg.), 160, 138-143 (1877).

nolds, 178 J. Moser, 179 and H. v. Helmholtz. 180 Tensions up to 3 atm were observed (Reynolds).

The other principal procedure was introduced by M. Berthelot 172 and is more suitable for detailed investigations. In this, the liquid is enclosed in a sealed tube which it nearly fills; by careful heating, the liquid is expanded until it completely fills the tube, exerting upon it a moderate pressure; then the temperature is slowly reduced. For a time, the liquid continues to fill the tube completely, but presently gives way with a snap, returning to the unstressed volume appropriate to the existing temperature. From the change in volume, Berthelot inferred that he had subjected water to a tension of 50 atm. The water was known to contain a small amount of dissolved air.

Improved apparatus employing Berthelot's procedure has been described and used by A. M. Worthington 181 and by J. Meyer 177 but the tensions did not exceed 34 atm. Meyer reported that the cooling curve is discontinuous at the instant of rupture.

H. H. Dixon 174 has subjected water containing dissolved air and fibers of wood to a tension of nearly 160 atm, using Berthelot's method. That is the highest recorded value that has been found.

From the adherence observed when flat, polished steel surfaces, very slightly wet with water, are wrung together, H. M. Budgett 181a concluded that the tension on the water at the time of rupture approached 60 atm. Before being placed together, the surfaces were wiped until they appeared dry to a casual observer. The water remaining on them formed isolated, microscopic drops. After wringing them together and then sliding them apart, it was found that the drops had been drawn out into thin parallel lines. It was estimated that the area of the ruptured surface of water did not exceed 1/10 of the complete area of the steel surface. By actual test, it was found that the surfaces would not adhere unless there was a minute quantity of liquid between them, and that their adherence was essentially the same in vacuo as in the free atmosphere.

O. Reynolds 171 observed that, when water is forced through a tube having at one point a short length of greatly constricted cross-section, a characteristic hissing is heard if the velocity exceeds a certain value. He described such experiments at the meeting of the British Association, in 1894, and attributed the hissing to the boiling of the water under the reduced pressure existing at the constriction. Were that the correct explanation, the hissing would begin at a very low velocity if the temperature of the water were nearly 100 °C. This has been tested by S. Skinner and F. Ent-

v. Helmholtz, H., Verh. deut. physik. Ges. (Berlin), 6, 16-18 (1887) = "Wiss. Abhand.," 3, 264-266, Leipzig, J. A. Barth, 1895.
 Worthington, A. M., Phil. Trans. (A), 183, 355-370 (1892).
 Budgett, H. M., Proc. Roy. Soc. London (A), 86, 25-35 (1912).

wistle. 182 They found that the velocity at which hissing begins is not zero at 100°, but over the range studied (12 to 99 °C) is essentially proportional to $(t_0 - t)$, indicating that it vanishes at, or near, the critical temperature (t_0) . From this they concluded that the hissing arises from collapse following a true rupture of the water, and that the tensile strength of water vanishes at a temperature near the critical.

J. Larmor 188 has shown that if the van der Waals equation continuously applies, the tensile strength of the liquid will vanish if the temperature equals or exceeds $(27/32)T_c$; which for water is 273 °C.

See also E. Askenasy, 184 G. A. Hulett 184a; and the compilation by T. F. Young and W. D. Harkins, 185

INTERNAL PRESSURE OF WATER

By the internal pressure of a liquid is meant the mean force of molecular attraction per cm2 required to maintain the molecules at their existing aver-

Table 81.—Internal Pressure of Water

 P_i = internal pressure; P = external pressure; v/v_{20} = ratio of the specific volume at t and P to that at 20 °C and 1 kg*/cm²; T °K = absolute temperature.

	P	= 1 kg*/cm ²			- Isometrics -	
Method•→ Ref.•→	State S	LtHt H	Visc `L		$P+P_i = T(\partial P/\partial t)$ TR	•
Rel.→		P_i		*	IK	P_i
0	11.66	12.69	72.6		$v/v_{20} = 2.11$	
10	11.60		74.6	525	1.692	9.688
20	11.50	12,49	71.3	575	2,348	7,632
30	11.44	12.32	68.0	625	2.819	3.701
40	11.35	12.17	57.2		$v/v_{20} = 2.13$	
50	11.24		49.7	525	1.640	9.190
60	11.13	11.84	43.8	575	2,278	7.852
70	11.02		39.0	625	2.762	3.858
80	10.92	11.45	35.2		$v/v_{20} = 2.18$	
90	10.81		31.9	525	1.508	8.312
100	10.71	11.04	25.9	575	2.082	6.948
120		10.60		625	2,565	5.235
140		10.11			$v/v_{20} = 2.25$	
160		9,63		525	1.368	7.202
180		9.14		575	1.876	6.264
		Other values		625	2.323	5.127

Unit of P and P = 1 megagram*'cm2 = 1000 kg*/cm2 = 967.8 atm. Temp. = t°C

[&]quot;Methods: LtHt = from the latent heat of vaporization. State = from an equation of state and the critical temperature. Visc = from the viscosity.

^b References:

Herz, W., Z. Elektrock., 32, 210-213 (1926). Lederer, E. L., Koll. Beih., 34, 270-338 (1932). Schuster, F., Z. anora. allgem. chem., 146, 299-304 (1925). Tammann, G., and Rühenbeck, A., Ann. d. Physik (5), 13, 63-79 (1932).

[°] From the effect of solutes upon the compressibility, P. G. Tait, 100 concluded that $P_i = 5.7$ at room temperature. P. Walden 107 computed for P_i at 100 °C the values: 11.0 from the latent heat, 8.4 from van der Waals' equation, and 4.4 from the surface tension.

age distances in opposition to the pressure arising from the thermal agitation of the molecules. Estimates of its value have been inferred in several ways from other types of data; it cannot be directly measured. Table 81.

VISCOSITY OF WATER

From a consideration of all pertinent data available in 1924, N. E. Dorsey 188 concluded that the values for the viscosity of water given in Tables 82, 85, and 86 are the best that can be derived from those data. They are the result of an entirely independent study of the recorded data, and involve many complete recomputations. In a forthcoming paper, the procedure followed will be described in some detail, and replies to certain criticisms of the conclusions reached will be given.

Those values for the range 0 to 100 °C at 1 atm differ by a few tenths of a per cent, usually in excess, from the corresponding ones published earlier by E. C. Bingham and R. F. Jackson, 189 and have been criticised by Bingham. 190 The greatest difference is nearly 0.5 per cent, and occurs near 17 °C (see Table 84). As the Bingham-Jackson values have been much used in the standardization of viscosimeters, they are here reproduced in Table 83, and compared with others in Table 84. In very many cases the precision of the measurements relative to water is such that the difference between these two sets of values is of no consequence; but in every case it is very desirable that both the temperature and the assumed viscosity of the water be explicitly stated so that future investigators may know how the results should be revised in order to correct them for any error that may have been discovered in the value of that assumed viscosity.

P. Leroux 191 ascribes an uncertainty of not over 1 in 200 to his elaborate determinations in the range 1.5 °C to 44.5 °C; but their variation with the temperature is quite different from that of the values obtained by others. It is believed that this discrepancy is due to errors in the temperatures, as the method by which he inferred the temperature of the water is not satisfactory, and the discrepancy is such as would exist if the recorded temperatures were, in each case, intermediate between the actual temperature of the water and that of the room, its difference from the actual temperature of the water increasing as that departs more and more from the temperature of the room, whether above or below. His values are omitted from this

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compilation. The only other sets of careful determinations of the viscosity of water that have come to the author's attention since 1924 are those given in sections II and III of Table 85. The determinations by G. A. Hawkins, H. L. Solberg, and A. A. Potter ¹⁹² are not satisfactory. The effect of eddies in the wake of the falling body has been ignored, and the temperature coefficients used for their instruments seem to be in error.

From observations by himself and by Beilby, L. Hawkes 198 inferred that the viscosity of water increases very greatly as the temperature is

Table 82.—Viscosity of Water: 0 °C to 109 °C

From compilation a by N. E. Dorsey. 188 See also text and Table 83.

The uncertainty in the tabulated values is probably of the order of 0.1 or 0.2 per cent between 0° and 40 °C, and of 0.5 to 1 per cent between 40 °C and 100 °C. Pressure = 1 atm. Temp. = $(t_1 + t_2)$ °C.

		U	nit of visc	cosity (η):	= 1 millipo	sise = 0.00	1 cgs unit.			
$t_2 \rightarrow t_1$	0°	1°	2°	3°	4°	5°	6°	7°	8°	9°
	42.04	45.00	46.74	16.10	7	4 . 40	44.50	44.00	40.05	40.40
0°	17.94	17.32	16.74	16.19	15.68	15.19	14.73	14.29	13.87	13.48
10	13.10	12.74	12.39	12.06	11.75	11.45	11.16	10.88	10.60	10.34
20	10.09	9.843	9.60_{8}	9.38_{0}	9.16_{1}	8.94	8.74_{6}	8.55_{1}	8.36	8.181
30	8.004	7.834	7.67_{0}	7.51,	7.357	7.20_{8}	7.064	6.92_{5}	6.79_{1}	6.66_{1}
40	6.536	6.415	6.29_{8}	6.184	6.07_{5}	5.970	5.86_{8}	5.770	5.675	5.58_{2}
50	5.49	5.40	5.32	5.24	5.15	5.07	4.99	4.92	4.84	4.77
60	4.70	4.63	4.56	4.50	4.43	4.37	4.31	4.24	4.19	4.13
70	4.07	4.02	3.96	3.91	3.86	3.81	3.76	3.71	3.66	3.62
80	3.57	3.53	3.48	3.44	3.40	3.36	3.32	3.28	3.24	3.20
90	3.17	3.13	3.10	3.06	3.03	2.99	2,96	2.93	2.90	2.87
100	2.84	2.82	2.79	2.76	2.73	2.70	2.67	2.64	2.62	2.59

"Based on the observations of:

Bingham, E. C., and White, G. F., Z. physik. Chem., 80, 670-686 (1912); Grotrian, O., Ann. d. Physik (Wicd.), 8, 529-554 (1879); Heydweiller, A., Idem, 59, 193-212 (1896); Hosking, R., Phil. Mag. (5), 49, 274-286 (1900); Idem (6), 7, 409-484 (1904); Idem, 17, 502-520 (1999); Idem, 18, 260-263 (1909); I. and Proc. Rov., Soc., N. S. Wales, 42, 34-56 (1908); Idem, 43, 34-38 (1909); Lyle, T. R., and Hosking, R., Phil. Mag. (6), 3, 487-498 (1902); Dr. Poiseuille, Mém. Savans Etraug. Inst. Paris, 9, 433-544 (1846); Compt. rend., 11, 961-967, 1041-1048 (1840); 12, 112-115 (1841); Idem, 15, 1167-1187 (1842); Slotte, K. F., Ann. d. Physik. (Wied.), 20, 257-267 (1883); Sprung, A., Idem (Pogg.), 159, 1-35 (1876); Thorpe, T. E., and Rodger, J. W., Phil. Trans. (A), 185, 397-710 (1894); and Washburn, E. W., and Williams, G. Y., J. Am. Chem. Soc., 35, 737-750 (1913).

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Table 83.—Viscosity of Water: 0 °C to 100 °C 189

Bingham and Jackson 189

Their values for the viscosity (η) are those defined by the formula $\eta^{-1} + 120 = 2.1482 \{(t - 8.435) + \sqrt{8078.4 + (t - 8.435)^2}\}$, in which the values of the constants were so determined as to fit a formula of this form to a certain set of mean values derived by them from the data available.

		Unit of	$\eta = 1 \text{ mill}$	ipoise = 0.	.001 cgs u	nit. Tem	$\mathbf{p}_{\cdot} = (t_1 + t_1)$	e)°C		
$t_2 \rightarrow t_1$	0	1	2	3	4	5	6	7	8	9
t_1					τ					
0	17.921	17.313	16.728	16.191	15.674	15.188	14.728	14.284	13.860	13.462
10	13.077	12.713	12.363	12.028	11.709	11.404	11.111	10.828	10.559	10.299
20	10.050	9.810	9.579	9.358	9.142	8.937	8.737	8.545	8.360	8.180
30	8.007	7.840	7.679	7.523	7.371	7.225	7.085	6.947	6.814	6.685
40	6.560	6.439	6.321	6.207	6.097	5.988	5.883	5.782	5.683	5.588
50	5.494	5.404	5.315	5.229	5.146	5.064	4.985	4.907	4.832	4.759
60	4.688	4.618	4.550	4.483	4.418	4.355	4.293	4.233	4.174	4.117
70	4.061	4.006	3.952	3.900	3.849	3.799	3.750	3.702	3.655	3.610
80	3.565	3.521	3.478	3,436	3.395	3.355	3.315	3.276	3.239	3.202
90	3.165	3,130	3.095	3.060	3.027	2.994	2.962	2.930	2.899	2.868
100	2.838									

Table 84.—Viscosity of Water: Comparison of Values

A comparison of (B) the Bingham and Jackson values of Table 83 with (D) those of Table 82 and with (M) Bingham and Jackson's "mean" of certain determinations. Those means seem to have served as the bases for the determination of the values of the 4 adjustable constants in their formula. The D values are here given as in *International Critical Tables*; the last figure has no significance.

			Unit	of $\eta = 1$ mic	cropoise = 1	0 d cgs unit.			
t	M - B	M	В	D	D-B	1	В	1)	D-B
0	- 34	17887	17921	17934	+ 13	10	13077	13097	+ 20
5	- 33	15155	15188	15188	0	11	12713	12733	+ 22
10	- 16	13061	13077	13097	+ 20	12	12363	12390	+ 27
15	+ 2	11406	11404	11447	+43	13	12028	12061	+ 33
20	- 4	10046	10050	10087	+ 37	14	11709	11748	+ 38
25	+4	8941	8937	8949	+ 12	15	11404	11447	+43
30	+ 12	8019	8007	8004	- 3	16	11111	11156	+45
35	- 20	7205	7225	7208	- 17	17	10828	10875	+47
40	- 27	6533	6560	6536	- 24	18	10559	10603	+44
45	- 30	5958	5988	5970	- 18	19	10299	10340	+41
50	+ 3	5497	5494	5492	- 2	20	10050	10087	+37
55	+ 8	5072	5064	5072	+ 8	21	9810	9843	+33
60	+ 13	4701	4688	4699	+ 11	22	9579	9608	+ 29
65	+ 4	4359	4355	4368	+ 13	23	9358	9380	+ 22
70	+ 1	4062	4061	4071	+10	24	9142	9161	+ 19
75	- 5	3794	3799	3806	+ 7	25	8937	8949	+ 12
80	- 9	3556	3565	3570	+ 5	26	8737	8746	+ 9
85	- 14	3341	3355	3357	+ 2	27	8545	8551	+ 6
90	- 19	3146	3165	3166	+ 1	28	8360	8363	+ 3
95	- 13	2981	2994	2994	0	29	8180	8181	+ 1
100	- 17	2821	2838	2839	+ 1	30	8007	8004	- 3

Table 85.—Viscosity of Saturated Water Below 0 °C and above 100 °C

Unit of $\eta = 1$ mp = 0.001 cgs unit; of $P_{\text{sat}} = 1$ kg*/cm² = 0.9678 atm. Temp. = t °C

I. Adapted from a compilation by N. E. Dorsey 188; see text. For t < 0 °C, values are by G. F. White and R. H. Twining, ²¹⁸ corrected and adjusted to accord with the values in Table 82. For t > 100 °C, values are from a table given by Heydweiller ²¹⁹; they are based on observations by M. de Haas 220 and have been so adjusted as to fit smoothly with the values tabulated by Thorpe and Rodgers (see Table 82, references) for temperatures below 100 °C. The three observations by de Haas (2.232 at 124.0 °C, 1.925 at 142.2 °C, and 1.805 at 153.0 °C) seem to be the only direct determinations of η at t > 100 °C that had been made before 1931 (cf. M. Jakob²²¹).

•	•							
£	Pant		η		1	1	Peat	η
- 2	0.0054	4	19.1		110	1	.46	2.56
- 4	0.004	6	20.5		120		.02	2.32
- 5	0.004		21.4		130		.75	2.12
- 6	0.004		22.0		140		.68	1.96
– 8	0.003	-	24.0		150		.85	1.84
- 10	0.003		26.0		160		.30	1.740
	0.000	•	20.0		100			1.72
II.	Adapted f	rom K	Sigwa	art. ²²²				
1	Punt		η		t	1	Pest	77
100	1.03		2.83		275	6	0.7	1.04
125	2.37		2.28		300		37.6	0.954
150	4.85		1.860		325		3.0	0.84
175	9.10		1.58		350		8.6	0.71
200	15.9		1.36°		360		0.4	0.63
225	26.0		1.23		370		4.7	0.53
250	40.6		1.13		374		25.2	0.378
200	40.0		1.10		014	24		0.576
III.	V. Shuga	ıyev ²²³	has rep	orted t	the followi	ng valu	ies:	
t	115.5	143.5	156.5	160.0	173.0	196.5	210.0	°C

1.82 mp ^a From the observed mobility of electrolytic ions, G. V. Hevesy ²²⁴ inferred the values: $\eta = 1.79$ at 156 °C, 1.21 at 218 °C, and 0.92 at 306 °C.

1.55

1.42

2.86

1.87

2.01

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Table 86.—Viscosity of Compressed Water

(For viscosity of saturated water $(P = P_{\text{sat}})$ see Table 85)

In Section I are given directly the values of the mean pressure coefficient of viscosity from 0 to P; from that and the value of η at 1 atm (essentially zero pressure) as given in Table 82, the viscosity under the pressure P can be computed. In Section II is given the only available set of data for compressed water above 100 °C.

Unit of
$$\eta = 1$$
 mp = 0.001 g/cm-sec = 0.001 cgs unit; of $P = 1$ kg*/cm² = 0.9678 atm; of $k = 10^{-6}$ per kg*/cm². Temp. = t °C.

I. Adapted from a compilation by N. E. Dorsey ¹⁸⁸ with the addition of computed values.

From their study of the viscosity of aqueous solutions, G. Tammann and H. Rabe ²²⁵ inferred that Bridgman's $(1926)^a$ values for water at $10\,^{\circ}$ C are in error, and set up expressions for the variation in the viscosity of water with the pressure, valid for $P \equiv 2000 \text{ kg*/cm}^2$. Those expressions are equivalent to the following: $10^6 (\eta - \eta_0)/\eta_0 P \equiv 10^6 k = -134.9 + 0.05778P$ at $0\,^{\circ}$ C, -37.63 + 0.02430P at $10\,^{\circ}$ C, +41.91 + 0.01054P at $30\,^{\circ}$ C, and +82.7 at $75\,^{\circ}$ C. Later, E. L. Lederer ²²⁶ set up the following equation for water:

$$\log_{10}(\eta/\eta_0) \equiv \log_{10}(1 + kP) = \frac{1.650P}{1000T} + \frac{1369P \cdot \log_{10}T}{10^8} - \frac{0.1300e^{-f}}{10^8}$$

where
$$f \equiv 10^{-6}(1350t^2) + \frac{691}{P}$$
; temperature = t° C, $T \equiv 273 + t^{\circ}$ C, and

unit of P=1 kg*/cm². They also tabulated the values of $\log_{10}(\eta/\eta_0)$ so computed for the values of P and T appearing in Bridgman's table. From these tabulated tables and from Tammann and Rabe's expressions were computed the several values of k here appearing under the heading "computed."

The observations of J. Sachs 227 and of E. Warburg and J. Sachs, 228 indicate that $10^{-6}k = -170$ per kg*/cm² at 20 °C for $P \ge 150$ kg*/cm². This does not agree with the other observations.

 $\eta=\eta_0(1+kP)$, where η and η_0 refer to the same temperature, but the first refers to the pressure P and the second to zero pressure, which may for our present purposes be taken as 1 atm. From the k here given and the value of η_0 as given in Table 82, η may be computed. For example, here we find for 30 °C and P=10000 kg*/cm², $k=+117\times10^{-6}$ per kg*/cm²; from Table 82 we find $\eta_0=8.00_4$ mp at 30 °C. Whence at 30 °C and 10 000 kg*/cm², $\eta=8.00_4(1+117\times10^{-6}\times10~000)=8.00_4(2.17)=17.4$ mp.

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	i		Unit of P	>= 1 kg*, cn	n²=0.9678	Juit of $P=1~{\rm kg^*,cm^2}=0.9678~{\rm atm;}$ of $k=10^{-6}~{\rm per}~{\rm kg^*/cm^2}.$	10-° per kg*	/cm². Tem	Temp. = t °C.	•			
t 17.94 0 17.94	10.3 12.99	xpermental 15 11.45	30 8.004	75 3.81	Ref.	0	10 Tammann	30 1 and Rabe	Computed 75	o l	10.3 Lea	30 derer	75
.8 -214° -128°	-200 ₆	-55 -63 -51	-474		ಜ ೦೦೦	-134 -129 -123 -118	-37.0 -35.2 -32.8 -30.4	+42.2 +43.0 +44.0 +45.1	+ 82.7 + 82.7 + 82.7 + 82.7	-84			
	-62	- 54 - 39 - 33	+49	+72	0 <u>m</u> 0 0 0	-112 -106 -100 -94	-27.9 -25.5 -23.0 -20.6	+46.1 +447.2 +48.2 +49.3	+ + 82.7 + 82.7 + 82.7	- 64	-72	+46	+ 9 4
2000 1000 - 79 2000 - 45 2000 - 45 4000 + 27.8 5000 + 44 6000 + 58 6000 + 58	- 46 - 29 - 16.0 - 16.0 + 433.2 + 443 + 452 + 60	0°.	++++++++++++++++++++++++++++++++++++++	++++++++++++++++++++++++++++++++++++++	Эщимимищищи	- 73. - 77.1 - 48.2 - 19.3 (+18.4)	$\begin{array}{c} -13.8 \\ -13.3 \\ -13.3 \\ -11.2 \\ +11.0 \\ (+35.3) \end{array}$	+51.4 +51.4 +52.4 +57.7 +63.0 (+73.5) (+94.6)	+82.7 +82.7 +82.7 (+82.7) (+82.7)	- 56 - 40 - 15 - 15 - 13 - 13 - 443 - 443 - 443	- 69 - 750 - 750 - 750 - 751 - 751 - 750 -	++++++++++++++++++++++++++++++++++++++	+ + + 97 + + 100 + + 1104 + + 1115 + + 127 + + 123 + + 133
310	362	20	1 1	30		Experimental (L. Hauser 50 +13	(L. Hauser 2	60	70	80	+	+13# +13# +82	516 40 40

Table 86 — (Continued)

II	From	K	Sigwart	222
TT	1.10111	1/	DIEWALL	

	•	3							
$t \xrightarrow{t} P_{\text{sat}} \rightarrow$	100 1 03	125 2 37	150 4 85	175 9 10	200 15 9	225 26 0	250 40 6	275 60 7	300 87 6
P					- η				
$P_{\mathtt{sat}}$	2 83	2 28	1 86	1 58	1 36	1 23	1 13	1 04	0 95
50	2 94	2 30	1 88	1 60	1 38	1 24	1 14		
100	3 01	2 44	1 94	1 64	1 41	1 26	1 15	1 05	0 96
200	3 24	2 53	2 03	1 70	1 46	1 28	1 19	1 08	0 99
300	3 43	2 67	2 14	1 76	1 51	1 32	1 21	1 11	1 02
$t \rightarrow$	325	350	360	370	374	400	410	430	450
$t \xrightarrow{t} P_{sat} \rightarrow$	123 0	168 6	190 4	214 7	225 2				
P					- η				
P_{sat}	0 84	0 71	0.63	0.53	0 378				
200	0 89	0 75	0 66	-					
300	0 93	0 82	0 76	071	0 69	0 40	0 33	0 29	0 28

a References

- Bridgman P W, Proc Am Acad Arts Sci 61, 57 99 (1926) Proc Nat Acad Sci, 11, 603 606 (1926) Cohen, R Ann d Physik (Wicd) 45, 666 684 (1892) Rontgen W C, Idem 22, 510 518 (1884) Others are given at head of tible

Table 87.—Viscosity of Sea-water 230

Salt content is s grams of salt per kg of sea-water, $\eta = viscosity$ of the sea-water, η_0 = viscosity of pure water at 0 °C η_t - viscosity of pure water at t °C as given in Table 82. The second half of the table has been computed from the first, by the present compiler

$s \rightarrow t$	5	10	$\frac{20}{1000\eta/\eta_0}$	30	40	5	10	$20 \\ 1000 \eta/\eta$	30	40
0	1009	1017	1032	1045	1059	1009	1017	1032	1045	1059
5	855	863	877	891	905	1010	1019	1036	1052	1069
10	738	745	758	772	785	1011	1020	1038	1057	1075
15	643	649	662	675	688	1008	1017	1037	1058	1078
20	568	574	586	599	611	1010	1021	1042	1065	1086
25	504	510	521	533	545	1010	1022	1044	1069	1093
30	454	460	470	481	491	1017	1031	1053	1078	1100

²²⁸ Shugayev, V Chem Abs 29, 2804 (1935) ← J Fxp Theo Phis (USSR) 4, 760 765 (1934)

At 9°C

[°]At 1°C

d At 23 °C

^{*} For the range 50 °C to 80 °C

²²⁴ Hevesy, G V Z Elektroch 27, 21 24 (1921)

²³⁸ Tammann, G and Rabe II 7 anorg allgem chem 168, 73 85 (1927)

²²⁶ Lederer E L, Koll Beth 34, 270 338 (1932)

²²⁷ Sachs, J. Diss Freiburg 1883

²²⁸ Warburg E Sachs J Ann d Physil (Wud) 22, 518 522 (1884)

²³⁹ Hauser, L, Ann d Phys (5) 5, 597 632 (1901)

²⁸⁰ Krümmel, O, and Ruppin, E. Wissensch Meeresunters (N F), 9, (Abt Kiel) 27 36 (1906) → Krümmel, O, "Handb d Ozeanog," Vol 1, 1907, Vol 2, 1911

reduced from -9 °C to -12 °C, and that water becomes a vitreous solid at -17 °C. But that is incorrect. It is incompatible with the data in Table 85, and is completely at variance with the compiler's observation that, to all appearances, water is essentially as mobile at -20 °C as it is at 0 °C. Furthermore L. Dufour ¹⁹⁴ stated that he had observed suspended drops of water to be fluid (flüssig) at -20 °C, and the observation of H. C. Sorby ¹⁹⁵ that the mobility of the clear liquid enclosed in small cavities in natural quartz is essentially the same at -20 °C as at room temperature has been quoted as evidence that water is quite mobile at -20 °C, and was so interpreted by him. The opinion, sometimes expressed, that such liquid inclusions are probably CO₂, not water, is contrary to the observations of Sir Humphry Davy ¹⁹⁶ (see p. 642).

Effect of Various Factors.

Magnetic Field.—W. König ¹⁹⁷ found that a magnetic field of 6300 to 7300 gauss transverse to the direction of flow of a solution of a paramagnetic salt produced no observable change in the viscosity of the solution. H. Dufour ¹⁹⁸ thought that he had observed that a transverse magnetic field decreased the viscosity of mercury, but as pointed out in the review, his observations are probably to be explained by the force exerted by the field on the flowing mercury. P. K. Raha and S. D. Chatterjee, ¹⁹⁹ using transverse fields up to 35000 gauss found no change for water, but definite changes for certain organic substances—for some an increase, for others a decrease. For a review of the subject, see M. Trautz and E. Fröschel. ²⁰⁰

Electric Field.—G. Pacher and L. Finazzi ²⁰¹ have found that an electrostatic field of nearly 27 kilovolt/cm transverse to the direction of flow of water produces no observable change in the viscosity. With other liquids the observed rate of flow in the field was very slightly different (not over 3 in 10,000) from that with no field, sometimes greater, sometimes less. They concluded that there is no true effect. This agrees with the observations of W. König ¹⁰⁷ but not with the conclusions reached by G. Quincke ²⁰² and by A. W. Duff, ²⁰³ which seem to be incorrect. (See criticism by Pacher and Finazzi ²⁰¹). For a solution of stearic acid in benzene O. Kimura ²⁰⁴ observed a marked increase in the viscosity. M. Trautz and E. Fröschel ²⁰⁵ have reviewed the subject.

Adjacent Solid.—(See also p. 512+). Some have suggested that when a viscous liquid is flowing over a solid there is a layer of the liquid of appreciable thickness, next to the solid, that remains at rest. But R. Bulkley ²⁰⁶ has found that in the case of oils, in which the effect should be especially pronounced, the thickness of such a stationary film does not exceed 0.02 μ to 0.03 μ (1 μ = 0.0001 cm). Similarly, F. P. Bowden ²⁰⁷ has been unable to find any evidence of surface forces acting at measurable distances; he could not work at distances smaller than 0.1 to 0.2 μ , but was certain that there is no such long-range effect (up to 50 μ) as some have reported. These results were confirmed by S. H. Bastow and F. P. Bowden.²⁰⁸ They

state: "No sign of induced rigidity was detected in liquids at a distance of 1000A $[0.1 \,\mu]$ from the surface even at temperatures near the freezing point. All the liquids investigated, except liquid crystals, were unable to withstand the slightest pressure without normal flow." Within experimental error, the viscosity of the thin film was the same as that of the liquid in bulk.

Although J. M. Macaulay ²⁰⁹ was undecided whether the high value (0.11 poise) that he computed from the rate at which water at 16 °C entered the gap between two parallel plates separated by 0.25 μ should be accepted as the actual value of the viscosity under those conditions, it seems most probable that it should not.

Dissolved Gas.—W. Ostwald and A. Genthe 210 have studied the effect of dissolved gas on the viscosity of water at 20 °C, and found as follows, η_0 being the viscosity of gas-free water:

Gas	N_2	O_2	CO ₂	CH.
η/η_0	1.017	0.990	1.007	0.998

Their report is lacking in detail. Ruppin found that the viscosity at 20 °C of air-saturated water is the same as that of gas-free water (see O. Krümmel and E. Ruppin.²¹¹

Relation between Viscosity and Other Properties.

Certain empirical and semitheoretical relations between the viscosity and other properties of the liquid have been proposed and discussed by D. B. Macleod,²¹² R. K. Sharma,²¹³ W. Herz,²¹⁴ D. Silverman and W. E. Roseveare,²¹⁵ and E. L. Lederer.²¹⁶

The theoretical expression $\eta = ATe^{-B/T}$ derived by J. Frenkel ²¹⁷ for the temperature variation of the viscosity of a liquid does not represent the observations on water.

30. Rigidity of Water (See also p. 189)

- F. Michaud ²³¹ has found that as the concentration of a jelly is progressively decreased, the rigidity becomes zero before the concentration does. This shows that the rigidity of water is zero. Like J. Colin, ²³² he quite disagrees with the conclusion of T. Schwedoff ²³³ regarding the rigidity of liquids.
- B. Derjaguin ²³⁴ has published the following values for the rigidity of very thin films of water:

Thickness of film	0.089	0.093	0.137	0.150	μ
Rigidity	1.9	1.7	0.04	0	10^8g*/cm^2

He suggests that the rigidity arises from chains of hundreds of oriented molecules extending from the solid boundaries into the liquid. His obser-

²³¹ Michaud, F., Ann. de Phys. (9), 19, 63-80 (1923).

vations are at variance with those of R. Bulkley²⁰⁶ on the viscous flow of oils, in which such chains might be expected and had been thought to exist. Bulkley found that adjacent to a solid wall there is no stationary film as much as $0.03 \,\mu$ in thickness. Behavior similar to that observed by Derjaguin would have resulted if the water had contained minute solid particles in suspension.

More recent work by F. P. Bowden and S. H. Bastow,²⁰⁸ B. Derjaguin,²³⁵ and J. M. Macauley ²⁰⁹ does not necessitate any change in the preceding statement.

31. Acoustic Data for Water

The greatest amount of vibratory energy that water at atmospheric pressure can transmit without cavitation is about 0.3 watt per cm^{2,286}

Velocity of Sound in Water.

The velocity of sound in water increases with the temperature to a maximum near 75 °C, and then decreases. In general, the velocity in other liquids decreases continuously.

Until 1927, the values obtained for the velocity of sound in water far from its boundaries, and for the variation of that velocity with the temperature, were quite discordant (cf. A. L. Foley ²³⁷). This was due in large part to the measurements having been made in vessels that were not large as compared with the wave-length of the sound employed, so that a large, complicated, and unsatsifactorily determined correction had to be applied to the observed velocity to eliminate the effect of the walls of the vessel.

In 1927, J. C. Hubbard and A. L. Loonis 238 published a very concordant set of preliminary data for frequencies (ν) of 200 to 400 kilocycles per second, and for temperatures of 5 to 35 °C. At such high frequencies, the waves are so short that containing vessels of moderate size produce no effect upon the observed velocity. This was followed the next year by a final report, in which the range was 0 to 40 °C. 239 And more recently, at this Bureau, very careful determinations have been made by C. R. Randall, 240 at $\nu = 750$ kilocycles/sec. In that work, the water was boiled in Pyrex glass immediately before being introduced into the apparatus, and extreme care was taken to ensure that the apparatus and contents had attained the temperature of the bath before measurements were made. The bath was thermostatically controlled to within \pm 0.02 °C. The uncertainty

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282 Colin, J., Compt. rend., 116, 1251-1253 (1893).
283 Schwedoff, T., J. de Phys. (2), 8, 341-359 (1889); 9, 34-46 (1890); (3), 1, 49-53 (1892).
284 Derjaguin, B., Z. Physik, 84, 657-670 (1933); Physik. Z. d. Sowj., 4, 431-432 (1933).
285 Derjaguin, B., Nature, 138, 330-331 (L) (1936).
286 Florisson, C., Bull. Soc. Belge Élect., 52, 165-170, 263-278, 339-348 (1936).
287 Foley, A. L., Int. Crit. Tables, 6, 464 (1929).
288 Hubbard, J. C., and Loomis, A. L., Nature, 120, 189 (1927).
289 Hubbard, J. C., and Loomis, A. L., Phil. Mag. (7), 5, 1177-1190 (1928).
280 Randall, C. R., Bur. Stand. J. Res., 8, 79-99 (RP402) (1932).
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241 Pooler, L. G., Phys. Rev. (2), 35, 832-847 (1930), superseding 31, 157 (A) (1928).

Table 88.—Velocity of Sound in Water

 V_o is the value defined by the empirical equation

$$V_o = 1404.4 + 4.8215t - 0.047562t^2 + 0.00013541t^3 = 1404.4 \times 10^{-1}$$

$$\left[1 + 3.4331_4 \left(\frac{t}{1000}\right) - 33.866_5 \left(\frac{t}{1000}\right)^2 + 96.419 \left(\frac{t}{1000}\right)^3\right]$$
 meters/sec;

it has a maximum at t = 74.2 °C. $V_{\rm obs} =$ observed velocity after application of a nominal correction for the finite size of the containing vessel (this correction is zero for the HL and the R values); $\Delta = V_{\rm obs} - V_c$, $\delta = \Delta/V_c$. In all cases the mean pressure is 1 atm, the boundaries are nominally at infinity, and the temperature is t °C. The R values are to be preferred (see text).

Unit of V_c and of $\Delta=1$ m/sec = 3.2808 ft/sec. Temp. = $t \circ C$

	V_{\circ}		_ Δ			- 1000δ -		_	Earlier	data -	
		Ra	HL^a	Pa `	Ra	HLa	Pa `	t	$V_{\mathfrak{o}}$	Δ	Ref.
0	1404.4	-0.9	+2.6		-0.6	+1.9		3.9	1422	-23	M
5	1427.3		+0.4			+0.3		7.6	1438	-29	M
10	1448.0	0	+0.8		0	+0.6		13.7	1462	-25	M M M B B
15	1466.5		+1.0			+0.7		25.2	1498	-41	M
20	1482.9	+0.2	+1.3		+0.1	+0.7		4.0	1423	-5^{b}	В
25	1496.30		+1.8	-10.3		+1.2	-6.9	21.5	1487	- 6b	В
30 35 40	1509.9	0	0.0	-11.2	0	0.0	-7.4	8.1	1440	- 5	CS
35	1520.7		-0.1			-0.1		13	1459	-18d	
40	1529.8	-0.3	+0.5	-11.5	-0.2	+0.3	-7.5	14	1463	-11	D
50	1543.5	0		-11.9	0		-7.7	18	1477	+12	D
60	1551.7	-0.2		-11.9	-0.1		-7.7	19	1480	- 19d	D
70	1555.3	0		-12.2	0		-7.8	19	1480	+38	D
75	1555.6			-11.4			-7.3	19	1480	+ 9	D
80	1555.0	-0.4			-0.3			31	1512	_ 7d	D
86	1553.4	-1.0			-0.6			19.5	1481	-19	С
								20	1483	-13	T

a References:

- B
- Bungetzianu, D., Bull. Soc. Roumaine des Sci., (Bucarest), 19, 1224-1246 (1910); 21, 208-267, 405-486 (1912); 22, 182-214 (1913). Especially the last. Cisman, A., J. de Phys. (6), 7, 345-352 (1926). Colladon, D., and Sturm, C., Mém. Sav. Etrano. Inst. Paris, 5, 267-347 (1838); Ann. de Chim. et Phys. (2), 36, 113-159, 225-257 (1827); Ann. d. Physik (Pogg.), 12, 39-76, 161-197 (1828); "Mémoire sur la compression des liquides et la vitesse de son dans l'eau," 1827, C. Schuchardt, Geneva, 1887. (All seem to refer to the same work.)
 Dörsing, K., Ann. d. Physik (4), 25, 227-251 (1909) Dies. Room. 1807. CS
- HL
- same work.)
 Dörsing, K., Ann. d. Physik (4), 25, 227-251 (1908) ← Diss., Bonn, 1907.
 Hubbard, J. C., and Loomis, A. L., Phil. Mag. (7), 5, 1177-1190 (1928); supersedes
 Nature, 120, 189 (1927).
 Jonesca, V., J. de Phys. (6), 5, 377-383 (1924).
 Martini, T., Atti R. Ist. Veneto (6), 4, Appendice (1886) (not included in index) →
 Beibl. Ann. d. Physik, 12, 566-568 (1888).
 Ref. 241.
 Ref. 240. J M
- ^b These two values differ slightly from the corresponding ones given in Foley's compilation and assigned to the same source. The one here given for 21.5 °C corresponds exactly to the value given by Bungetzianu; that for 4 °C was computed by the compiler from the data given by the observer.
- ^o L. Bergmann, ²⁴⁷ using frequencies of 4.5 to 13 megacycles/sec and inferring the length of the waves from their observed diffraction of light, obtained for the velocity at 25 °C the very low value, 1465 m/sec. But S. Parthasarathy, 248 using a similar method and a frequency of 7.32 megacycles/sec, found 1494 m/sec at 24 °C.
- ⁴ These three were obtained in the same glass tube and at the same frequency; the others by D were obtained in other tubes and, in part, at other frequencies.

²⁴² Boyle, R. W., Lehmann, J. F., and Morgan, S. C., Trans. Roy. Soc. Canada, III (3), 22, 371-378 (1928).

²⁴⁸ Boyle, R. W., and Taylor, G. B., Idem, 21, 79-83 (1927), superseding 19, 197-203 (1925).

²⁴⁴ Biquard, P., Compt. rend., 188, 1230-1232 (1929).

Table 89.—Velocity of Sound in Natural Waters

(See also Table 90)

Excepting the references to Lübcke and to Dorsey, the following has been taken with slight changes in form from the compilation of A. L. Folev.287

In all oceans the average vertical velocity for depths of 3.5 to 8.0 km (2.2 to 5 miles) is 1528 to 1529 meters/second; for lesser depths it is less. In general, the horizontal velocity increases by 0.2 per cent per 1 °C increase in temperature, 0.11 per cent per 100 meters increase in depth, and 0.1 per cent per 1 per cent increase in salinity (total salts).²⁴⁹ A. B. Wood and H. E. Browne ²⁵⁰ represent the data (6 to 17 °C, salinities near 3.5 per cent) obtained by A. B. Wood, H. E. Browne, and C. Cochrane ²⁵¹ by means of the equation $V = 4626 + 13.8t - 0.12t^2 +$ 3.73s ft/sec = $1410.0 + 5.21t - 0.036t^2 + 1.137s$ m/sec, where the salinity is s parts per 1000 and the temperature is t °C. The practical application of acoustics to coastal and oceanic surveying has been discussed by H. G. Dorsey, 252 who gives data indicating that the velocity is independent of the intensity of the source.

Unit of V=1 meter/sec; of depth (D)=1 meter; of salinity (s)=1 per cent by weight. Temp. =t °C.

	Ocea	an: Horizontal	Velocity.		
Place	D	s	t	\boldsymbol{v}	Ref.
Open ocean	13	36	14.5	1503.5	M
Block Island Sound, N. Y.	30	3.35	3.0	1453.3	S
Long Island Sound, N. Y.	30		13	1492.3	E
Isle of Wight		3.51	6	1474	WBC
Isle of Wight		3.52	7	1478	WBC
Isle of Wight		3.5	16.95	1511	WBC

	Uccan: Vertical	velocity	
Place	D	V	Ref.
N. Atlantic	1288	1520	HS
Carribean Sea	338	1478	HS
Carribean Sea	1771	1486	HS
Pacific	1185	1505	HS
Pacific	2962	1493	HS
All oceans	>3500	1528	HS
		- Fresh Water -	
Water	t	V	Ref.
Tales Comores	0 1	1/25	CC

Ref.
CS
W
W
W
W

a References:

CS See Table 88, references.

E Eckhardt, E. A., Phys. Rev. (2), 24, 452-455 (1924).

HS Heck, N. H., and Service, J. C., U. S. Coast and Geod. Sur., Spec. Publ. 108 (1924).

M Marti, Compt. rend., 169, 281-282 (1919).

S Stephenson, E. B., Phys. Rev. (2), 21, 181-185 (1923).

W Wertheim, G., Ann. de Chim. et Phys. (3), 23, 434-475 (1848) = Ann. d. Physik (Pogg.), 77, 427-445, 544-571 (1849).

WBC Ref. 251.

Density at 14.9 °C was 1.0245 g/cm³.

Table 90.—Velocity of Sound in Sea-water

(See also Table 89)

Adapted from the detailed practical tables by N. H. Heck and J. H. Service 253 based upon the very extensive tables of V. Bjerknes et al. 254

The following values of the velocity (V) at 0 °C and various depths, salinity 35 g/kg, have been taken directly from the tables of Heck and Service. The values they give for other salinities (s = 31 to 37 g/kg) and temperatures $(t = 0 \text{ to } t_m \, ^{\circ}\text{C})$ may be reproduced very closely by means of the formula

$$V_{s,t,d} = V_{35,0,d} + 2.39t - 0.028t^2 + \{0.83 - 18.0(10^{-6})d + 0.0075t\}(s - 35)$$

the units being those named below.

Heck and Service compute the velocity for each successive layer of 200 fathoms, inferring the distribution of temperature and salinity from observations at three depths, i.e., surface, 200 fathoms, and bottom, and average these velocities to obtain the mean velocity (V_m) . They find in actual practice that this mean velocity differs from that computed from the measured depth and the observed time required for sound to pass to the bottom and back, by an average of about 0.2 per cent (about 3 m/sec), the probable error for a single determination being 6 to 8 times as great, and single determinations differing from V_m by 3.5 per cent (52.5 m/sec). (In his compilation,²³⁷ A. L. Foley seems to have had in mind the probable error of a single determination when he stated that values computed by the method of Heck and Service may differ by 20 m/sec from the actual value). Such differences include the errors inherent in the method of echo-sounding as well as those arising from an attempt to infer the distribution of temperature and salinity from observations at only three depths.

	Unit of d	$= d_1 + d_2 =$	= 1 fatho	m = 6 ft. =	182.88	cm; of V	= 1 fath	om/sec =	1.829 m/s	sec.
$d_1 - d_2$	100	300	500 - V 25,0,d	700	900	100	300	500 t_m	700	900
0	793	796	799	804	806	22	22	22	20	12
1000	809	813	816	820	825	8	6	4	4	3
2000		831	833	836	839	3	3	3	3	2
3000		848	850	855	857	2	2	2	2	2
4000	861	863	866	870		2	2	2	2	
$d_1 - d_2$	100	300	500	700 V 15,0,d	900 (Unit of	V = 1 m/s	1300 sec)	1500	1700	1900
0	•	1456	1461	1470	1474	1479	1487	1492	1500	1509
2000	1510	1520	1523	1529	1534	1544	1551	1554	1564	1567
4000	1574	1578	1584	1591						

²⁴⁵ Špakovskij, B., Compt. rend. Acad. Sci. URSS (N. S.), 1934₈, 591-594 (1934).

²⁴⁶ Schaffs, W., Z. Physik, 105, 658-675 (1937).

²⁴⁷ Bergmann, L., Physik. Z., 34, 761-764 (1933).

²⁴⁸ Parthasarathy, S., Proc. Indian Acad. Sci. (A), 2, 497-511 (1935).

²⁴⁹ Lübcke, E., Z. techn. Physik, 10, 386-388 (1929).

²⁵⁰ Wood, A. B., and Browne, H. E., Proc. Phys. Soc. (London), 35, 183-193 (1923).

²⁵¹ Wood, A. B., Browne, H. E., and Cochrane, C., Proc. Roy. Soc. (London) (A), 103, 284-303 (1923).

in the values obtained is believed to be distinctly less than 0.1 per cent. They are the values to be preferred.

L. G. Pooler ²⁴¹ has measured the velocity at 25 to 75 °C and $\nu = 1269$ to 2715 cycles per second, using a recently developed formula for correcting for the effect of the walls of the vessel. His values lie 0.7 to 0.8 per cent below those of Randall and of Hubbard and Loomis (see Table 88).

To facilitate comparison of the several sets of data, an empirical formula (1) of arbitrary form, but reproducing Randall's values at 10, 30, 50, and 70 °C, was set up. The values so determined are designated as V_o .

$$V_c = 1404.4 + 4.8215t - 0.047562t^2 + 0.00013541t^3 \text{ m/sec}$$
 (1)

The excess of the reported velocity (V_{obs}) above V_c is given in Table 88 for each of a number of determinations.

The observations of R. W. Boyle, J. F. Lehmann, and S. C. Morgan 242 at 80 kc/sec, of Hubbard and Loomis (200 to 400 kc/sec), of R. W. Boyle and G. B. Taylor ²⁴³ (29 to 570 kc/sec), of Randall (750 kc/sec), and of P. Biguard ²⁴⁴ (1360 kc/sec) indicate that the velocity varies little, if at all, with the frequency. In fact, there is no certain reported evidence that the velocity in an unbounded volume of water changes at all as the frequency is varied from 100 cycles/sec to 1.4 megacycles/sec. The smaller values reported for the lower frequencies (see Table 88, columns headed "P" and "Earlier data") probably arise from the unsatisfactory nature of the correction which is necessitated in such cases by the presence of the walls of the vessel.

More recently, B. Špakovskij, 246 using values of v up to 1000 kc/sec, has concluded that up to that frequency, at least, the velocity is constant within his experimental error (about 1 per cent), and W. Schaffs 246 using $\nu = 16381$ kc/sec, found the velocity to be 1467 m/sec at 17 °C. This is only 0.3 per cent smaller than the value defined by formula (1). Whence it seems probable that the velocity is independent of the frequency, at least up to 16 megacycles/sec.

Acoustic Resistivity of Water.

The acoustic resistivity of a material is defined as the amount by which the r.m.s. pressure in a plane sound wave must exceed the static pressure in order to confer upon the medium a unit r.m.s. velocity. (r.m.s. = square root of the mean square.) It is equal to $\sqrt{E_{\rho}} = V_{\rho}$, where E = bulkmodulus of the material, ρ is its density, and V = velocity of sound = $\sqrt{E/\rho}$. For water at usual temperatures, ρ is essentially unity, and the acoustic resistivity is numerically equal to the velocity of sound. If the velocity is V meters per second, the resistivity is V gram/mm²-sec. In each, the modulus is that corresponding to the conditions existing during

²⁵³ Dorsey, H. G., J. Acoust. Soc. Amer., 3, 428-442 (1932).

²⁸⁸ Heck, N. H., and Service, J. H., U. S. Coast and Geod. Sur., Spec. Publ. 108 (1924).
288 Bjerknes, V., et al., "Dynamic Meteorology and Hydrography," Carnegie Inst. of Washington Publ. 88 (1910).

the passage of the wave, which approximate those characteristic of adiabatic compression and expansion.255

H. G. Dorsey 252 remarks that the increase in the acoustic resistance of sea-water with increase in temperature may not arise solely from the change in temperature, but may be due in part to an increase in the amount of suspended matter. He states that water churned up by the propeller of a ship absorbs sound completely, and that the reflectivity of ocean bottoms decreases in the order: soft mud (best), hard sand, broken coral, sea-grass (poorest).

In a more recent paper ²⁵⁶ he writes (p. 299): "The assumption that the acoustical impedance of warm water is greater than cold has not been disproved and work in the warm water of the Gulf of Mexico tends to confirm the assumption."

Absorption of Sound by Water.

At audio frequencies the absorption of sound by water is small, and is determined by the viscosity and thermal conductivity. But at high—ultrasonic—frequencies it is much greater and seems to involve something of the nature of molecular resonance. Furthermore, at those high frequencies the radiation has a degassing effect; and this absorbs additional energy if the water is not gas-free. (See C. Sörensen.²⁵⁷)

Sörensen 257 has found that, although the absorption is greater if the water contains gas, the heating of the water by the radiation is the same as if the water were gas-free, the additional absorption arising from the work required to drive out the gas. He found as follows: for $\nu = 194,380$, and 530 kc/sec, the work expended in removing the gas was, respectively, 51.2, 72.6, and 87.4 kilowatts per cm³ of gas removed. The rate of removal was not constant, but steadily decreased as the water became more and more nearly gas-free.

He stated that the absorption passes through a maximum at some frequency below 194 kc/sec. H. Oyama 258 has stated that the heating of the water is a maximum at about 700 to 800 kc/sec, which seems to be entirely incompatible with the values in Table 91.

Claeys found the absorption to be greater in narrow tubes than in wider ones, and has suggested that the difference is associated with the presence of convection currents 259; but Sörensen 260 found the absorption to be independent of the diameter of the tube.

Sörensen 260 has found that the coefficient of absorption (k, see Table 91) decreases as the temperature rises, but that this decrease is less rapid

²⁵⁵ Sabine, P. E., Int. Crit. Tables, 6, 459 (1929).

²⁵⁶ Dorsey, H. G., J. Acoust. Soc. Amer., 7, 286-299 (1936).

²⁶⁷ Sörensen, C., Ann. d. Physik (5), 26, 121-137 (1936); 27, 70-74 (1936). 268 Oyama, H., Sci. Abs. (A), 39, 292 (1936) ← J. Inst. Elec. Eng. (Japan), 55, 985-989 (1935).

²⁵⁹ Claeys, J., and Sack, H., Acad. Roy. de Belg., Bull. Cl. Sci. (5), 23, 659-671 (1937).

²⁰⁰ Sörensen, C., Ann. d. Physik (5), 27, 70-74 (1936).

than that of $\eta/\rho v^8$, where η is the viscosity, ρ the density, and v the velocity of sound. The variation is not linear in the temperature, but near room temperatures dk/dt is quite close to -0.00024 cm⁻¹ per 1 °C, the frequency being between 200 and 1000 kc/sec. E. Baumgardt, 261 on the other hand, concluded that k is proportional to $n/\rho v^3$, and reported the following values. all for $\nu = 7958 \text{ kc/sec}$:

t	18.6	22.2	22.5	31.2	39.5 °C
100k	3.46	3.06	2.98	2.42	2.08 cm ⁻¹

That k is not always proportional to v^2 was pointed out by P. Biquard, ²⁶² and is obvious from the data in Table 91. With toluene, Biquard 263 observed a lateral scattering of the radiation.

Claeys, Errera, and Sack ²⁶⁴ have suggested that the increased absorption at high frequencies may arise in part from a kind of hysteresis in the adiabatic compressibility of the water.

Table 91.—Absorption of Ultrasonic Radiation by Water

The coefficient k is that defined by the relation $I = I_0 e^{-k\omega}$ where $I_0 - I$ is the reduction in the intensity of a plane wave while traveling a distance x. Data have been published both in terms of k and of the corresponding exponent for the reduction in amplitude, which is only half as great as k, and it is not always easy to determine to which they refer.

Unit of $\nu = 10^6$ cycles/sec; of k = 1 cm⁻¹; of $k/\nu^2 = 10^{-14}$ sec²/cm. Room temp.

• 0.	10 0,010,000,010 100,000		Tito out tour by
ν	k	k/>2	Ref.
0.194	0.017	45	S
0.380	0.015	10	S
0.530	0.014	5.0	8 S 8 S
0.950	0.011	1.22	S
1.44	0.00135	0.065	CES
2.03	0.00330	0.080	CES
2.79	0.0030	0.038	\mathbf{F}
4.77	0.0121	0.053	CES
7.55	0.035	0.062	$\mathbf{B}_{\mathbf{i}}$
7.96	0.0346	0.055	Ba
7.97	0.040	0.064	Bi
8.37	0.027	0.038	F
11.14	0.066	0.054	CES
54	1.39	0.048	Bär
69	2.28	0.048	Bär
83	3.52	0.051	Bär

* References:

Ba Bär Bi

CES

Baumgardt, E. 201 Bar, R., Helv. Phys. Acta, 10, 332-337 (1937). Biquard. P. 200, 2005 Claeys, J., Errera, J., and Sack, H. 204 Fox, F. E., Phys. Rev. (2), 52, 973-981 (1937). Sörensen, C., Ann. d. Physik (5), 26, 121-137 (1936) = Diss., Greifswalder, 1935.

²⁶¹ Baumgardt, E., Compt. rend., 202, 203-204 (1936).

²⁰² Biquard, P., Ann. d. Phys. (11), 6, 195-304 (1936).
203 Biquard, P., Compt. rend., 202, 117-119 (1936).

²⁰⁴ Claeys, J., Errera, J., and Sack, H., Idem, 202, 1493-1494 (1936).

⁹⁰⁸ Biquard, P., Compt. rend., 193, 226-229 (1931).

32. Pressure-Volume-Temperature Associations for Water (For saturated water, see Section 88)

Water under the pressure of its pure saturated vapor is called saturated water; that under a higher pressure has been called compressed water; that under a lower pressure may be called dilated water. Above the critical temperature the substance will be classed as compressed water if the specific volume is less than that (3.1 cm³/g) at the critical point; as dilated vapor if the specific volume is greater than at the critical point. These terms are to be so understood wherever they occur in this compilation.

Uniformity of Water.

Until the discovery of the isotopes of hydrogen and oxygen—that is, until very recently-water as commonly purified by careful distillation was (To p. 202)

Table 92.—Effect of Dissolved Air on the Density of Water (See p. 251)

 ρ = density of air-free water under a pressure of 1 atm; ρ_a = density of water saturated with air at a pressure of 1 atm.

	Unit of ρ and of $\rho_a =$	1 g/cm³.	Temp. = t °C.		
t	$10^{6}(\rho-\rho_a)$ Observer ^a	t	$10^6(\rho-\rho_a)$	t	$10^6(\rho-\rho_a)$
5° to 8 °C	3.0 Chappuis		Marek		Marek
15.6	1.89 Frivold	6	3.3	13	2.7
0	2.5 Marek	7	3.4	14	2.5
1	2.7 Marek	8	3.4	15	2.2
2	2.9 Marek	9	3.3	16	1.9
3	3.1 Marek	10	3.2	17	1.6
4	32 Marek	11	3.1	18	1.2
5	3.3 Marek	12	2.9	20	0.4

^a See text for comments on Marek's work and references to his and to Chappuis' papers. Frivold.250

²⁸⁶ Lamb, A. B., and Lee, R. E., J. Am. Chem. Soc., 35, 1667-1693 (1681) (1913).

²⁰⁷ Hall, N. F., and Jones, T. O., J. Am. Chem. Soc., 58, 1915-1919 (1936); Gabbard, J. L., and Dole, M., Idem, 59, 181-185 (1937).

²⁰⁸ Christiansen, W. H., Crabtree, R. W., and Laby, T. H., Nature, 135, 870 (L) (1935).

²⁰⁰ Mendelejev, J., Compt. rend. Acad. Sci. URSS, 8, 105-108 (1935₃).

²⁷⁰ Dole, M., and Wiener, B. Z., Science (N. S.), 81, 45 (1935).

²⁷¹ Peel, J. B., Robinson, P. L., and Smith, H. C., Nature, 120, 514-515 (1927).

²⁷² Stott, V., and Bigg, P. H., Int. Crit. Tables, 3, 24-26 (1928).

²⁷⁸ Chappuis, P., Trav. et Mém. Bur. Int. Poids et Mes., 13, D1-D40 (1907).

²⁷⁴ Thiesen, M., Scheel, K., and Diesselhorst, H., Wiss. Abh. Phys.-Techn. Reichs., 3, 1-70 (1900).

²⁷⁵ Tilton, L. W., and Taylor, J. K., J. Res. Nat. Bur. Stand., 18, 205-214 (RP971) (1937).

²⁷⁶ Bridgman, P. W., Int. Crit. Tables, 3, 40 (1928); as corrected in accordance with the errata published in Vol. 7.

²⁷⁷ Bridgman, P. W., Proc. Am. Acad. Arts Sci., 47, 439-558 (1912); 48, 307-362 (1913).

²⁷⁸ Bridgman, P. W., J. Chem'l Phys., 3, 597-605 (1935).

Table 93.—Density of Compressed Water at a Pressure of 1 Atmosphere

(For sea-water see Table 108.)

Density = ρ , temperature = $(t_1 + t_2) = t$ °C. In the second column the complete value of ρ is given or indicated; in the following columns, only the last three or four digits, the preceding digits being understood to be those in the left-hand section of the second column, either on or above the line, unless there is a line over the first of the tabulated digits, in which case the immediately preceding digit will differ by one unit from that just specified. For example, Mohler finds $\rho = 0.997292$ at -12 °C and 0.996931 at -13 °C.

Unit of $\rho = 1$ gram per milliliter = 0.999973 g/cm³. Temp. = $(t_1 + t_2) = t$ °C.

I. J. F. Mohler.²⁹⁰

II. C. Despretz.²⁸⁰ Not given in *International Critical Tables*; he studied the expansion of water between -9 and +100 °C.

t
 0
 -1
 -2
 -3
 -4
 -5
 -6
 -7
 -8
 -9

$$\rho$$
 0.999 873
 786
 692
 578
 438
 302
 082
 $\overline{8}48$
 $\overline{6}28$
 $\overline{3}72$

 t
 0
 1
 2
 3
 4
 5
 6
 7
 8
 9

 ρ
 0.999 873
 925
 967
 992
 $\overline{0}00$
 992
 969
 929
 878
 812

Revised Chappuis table.²⁷⁵ See text. In the subsidiary columns C and T to the right of the ρ values in columns $t_1 = 0.0$ and $t_1 = 0.5$ are given the amounts, in units of the last place tabulated, by which each of the corresponding values for $t_1 = 0.0$ and $t_1 = 0.5$ in the tables published by Chappuis and by Thiesen, Scheel, and Diesselhorst, respectively, exceeds that here tabulated. Over the intermediate 0.5 °C range these differences may be linearly interpolated. Thus both of those tables may be recovered from this; and so may be the one in the International Critical Tables, that being merely the average of the other two. For example, at 17.3 °C this table gives $\rho = 0.9987515$, the C value is smaller than this by 10, the T value by 27, and the I.C.T. by 18 units in the last place, making those values 0.9987505, 0.9987488, and 0.9987497 respectively; from the tables themselves one finds exactly these same values. Under Δ is given the average increase in ρ per 0.1 °C increase in t for the one degree range covered by the line in which the value stands, the unit of Δ being that of the last tabulated digit of p.

0.0 c T C T	0.99	0.0														
C T 8676 +5 0 8743 8808 8871 8933 +3 +1 9051 9107 9161 9214 9265 +2 +1 9314 9362 9407 9451 9493 +1 +2 9573 9610 9645 9678 +1 +2 9710 9740 9760 9821 0 9886 9903 9907 9909 9922 0 0 9937 9950 9962 9962 9968 9993 9997 9997 9000 0 9937 9950 9962 9962 9981 9884 9993 9993 9997 9010 0 0 9988 9962 9966 9970 9760 9770 9742 9713 9010 0 9984 9864 9843 9826 9970 9749 9749 9749 9749 9749 9748 9749 9749 97	0.99				0.1	0.2	0.3	9.4	0.5			9.0	0.7	0.0	6.0	+0.1°
8676 + 5 0 8743 8808 8871 8933 8993 + 3 + 1 9051 9101 9101 9101 9101 9101 9101 9101 9101 9101 9101 9101 9101 9101 9101 9102 9602 9902 9102 981 961 966 9	0.90		O	T						ပ	T					1
9265 + + + 1 9314 9362 9407 9451 9493 + 1 + 2 9534 9573 9610 9645 9678 + 1 + 2 9710 9740 9760 9821 0 + 1 9844 9860 9886 9905 9922 0 0 9377 9950 9962 9972 9981 0 9988 9993 9997 9996 9000 0 0 9999 9996 9992 9986 9972 9981 9986 9993 9997 9999 9996 9996 9996 9997 9998 9993 9997 9998 9993 9997 9998 9993 9997 9998 9993 9997 9998 9993 9997 9998 9997 9998 9997 9999 9996 9996 9998 9997 9999 9994 9884 9864 9848 9864 9848 9869 9993 9997 9898 9997 9896	1.00		+5	0	8743	8808	8871	8933	8993	+3	7	9051	9107	9161	9214	+58.9
9678 +1 +2 9710 9740 9769 9821 0 +1 9844 9866 9886 9905 9922 0 0 9937 9950 9952 9972 9981 0 0 9988 9993 9997 9999 9000 0 9937 9966 9962 9972 9981 0 9988 9996 9996 9996 9996 9996 9997 9998 9996 9997 9998 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9997 9998 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9997 9998 9998 9998 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996 9996	1.00	-	+2	7	9314	9362	9407	9451	9493	+1	+2	9534	9573	9610	9645	+41.3
9922 0 0 9937 9950 9962 9972 9981 0 0 9988 9993 9997 9999 99900 9900 9900 9900 990	1.00	8496	+	+2	9710	9740	6916	9646	9821	0	+1	9844	9986	9886	9905	+24.4
0000 0 0 9999 9996 9992 9986 9979 0 -1 9970 9969 9996 9999 9984 9885 9889 9889 9889 9889 9889 9889 9889 9889 9889 9889 9889 9889	1.00	9922	0	0	9937	9950	9965	9972	9981	0	0	8866	9993	1666	6666	+7.8
9919 0 -1 9902 9884 9864 9843 9820 -1 2 9706 9770 9742 9713 9683 -1 -3 9651 9618 9583 9546 9508 -1 -3 9469 9428 9386 9342 9297 -1 -4 9250 9202 9153 9102 9049 0 -4 8995 8940 8883 8825 8765 -1 -6 8704 8642 8578 8513 8446 -1 7 8796 8238 8166 8092 -1 -6 8704 8642 8578 8513 8446 -1 -7 8378 8166 8338 8166 8338 8166 8883 8825 8238 8166 8374 8518 7784 7704 0 -1 7 8799 8238 8166 8238 8166 82374 8236 8136 8136	000	000000	0	0	9999	9666	9992	9866	9979	0	-1	0266	<u>0</u> 966	9948	9934	-8.1
9683 -1 -3 9651 9618 9536 -1 -3 9469 9428 9386 9342 9297 -1 -4 9250 9202 9153 9102 9049 0 -4 8995 8940 8883 8855 8765 -1 -6 8704 8642 8578 8513 8446 -1 7336 9490 8833 8825 8092 -1 -8 8017 7941 7863 7784 7704 0 -10 7622 7539 7454 7368 7281 +1 -10 7193 7103 7012 6919 6825 +1 -1 8399 8238 8166 5261 -13 6027 5922 5815 -10 -14 5706 5397 5486 5374 4059 -19 -18 3932 3803 3674 3543 3411 -20 -19 4534 4432	``	99 9919	0	1	9902	9884	9864	9843	9820	-	-2	9616	9770	9742	9713	-23.6
9297 -1 -4 9250 9202 9153 9102 9049 0 -4 8995 8940 8883 8825 8765 -1 -6 8704 8642 8578 8513 8446 -1 7 8378 8309 8238 8166 8092 -1 -8 8017 7941 7863 7784 7704 0 10 7622 7539 7454 7368 7281 +1 -10 7193 7103 7012 6919 6825 +1 -1 7620 7539 7454 7368 5261 -13 6027 5922 5815 -10 -14 5706 5597 5486 5374 4059 -19 -18 3932 3803 3674 3543 3411 -20 -19 4534 4432 4430 4184 4059 -19 -18 3932 3803 3674 3543 3411		9683	7	-3	9651	9618	9583	9546	9508	11	13	9469	9428	9386	9342	-38.6
8765 -1 -6 8704 8642 8578 8513 8446 -1 7 8378 8309 8238 8166 8092 -1 -8 8017 7941 7863 7784 7704 0 10 7622 7539 7454 7368 7281 +1 -10 7193 7103 7012 6919 6825 +1 -1 6730 6634 6536 6437 6336 -5 -12 6234 6131 6027 5922 5815 -10 -14 5706 5597 5486 5374 -18 5261 -13 6027 5922 5815 -10 -14 5706 5597 5486 5374 -18 4059 -19 -18 3932 3803 3674 3543 3411 -20 -19 1379 4134 4130 4184 -19 4132 4139 4184 -19 4131 4136		9297	7	4	9250	9202	9153	9102	9049	0	4-	8995	8940	8883	8825	-53.2
8092 -1 -8 8017 7941 7863 7784 7704 0 -10 7622 7539 7454 7368 7281 +1 -10 7193 7103 7012 6919 6825 +1 -11 6730 6634 6536 6437 6336 -5 -12 6234 6131 6027 5922 5815 -10 -14 5706 5597 5486 5374 -586 5374 -586 5374 -586 5374 -586 5374 -586 5374 -586 5374 -586 5374 -586 5374 -586 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 5374 -588 53		8765	-1	9-	8704	8642	8578	8513	8446	-	1-1	8378	8309	8238	8166	-67.3
7281 +1 - 10 7193 7103 7012 6919 6825 +1 - 11 6730 6634 6536 6437 6336 -5 - 12 6234 6131 6027 5922 5815 -10 - 14 5706 5597 5486 5374 - 5261 -13 - 15 5146 5030 4913 4795 4675 -15 - 16 4524 4432 4309 4184 4059 -19 - 18 3932 3803 3674 3543 3411 -20 - 19 3278 3143 3007 2870 2732 -20 - 19 2593 2453 2311 2168 2024 -21 - 21 1879 1732 1584 1436 2732 -20 - 22 1134 0982 0828 0674 0518 -19 - 23 0360 0202 0043 982 9721 -16 - 24 9558 9394 9229 9062 8895 -14 - 25 8726 857 8386 8214		8092	-1	80	8017	7941	7863	7784	7704	0	- 10	7622	7539	7454	7368	-81.1
6336 -5 -12 6234 6131 6027 5922 5815 -10 -14 5706 5597 5486 5374 5261 -13 -15 5146 5030 4913 4795 4675 -15 -16 4554 4432 4309 4184 4059 -19 -18 3932 3803 3674 3543 3411 -20 -19 3778 3143 3007 2870 2732 -20 -19 2593 2453 2311 2168 2024 -21 -21 1879 1732 1584 1436 2732 -20 -19 2593 2453 2311 2168 2024 -21 -21 1879 1732 1584 1436 2721 -10 -24 9558 0922 9062 8895 -14 -25 8726 8577 8386 8214 2724 -12 -27 7867 7691 7515 7337 7158 -8 -27 6979 6798 6616 6433 2748 -1		7281	+1	-10	7193	7103	7012	6169	6825	+	-11	6730	6634	6536	6437	-94.5
5261 -13 - 15 5146 5030 4913 4795 4675 -15 - 16 4554 4432 4309 4184 4059 -19 - 18 3932 3803 3674 3543 3411 -20 - 19 3778 3143 3007 2870 2732 -20 - 19 2593 2453 2311 2168 2024 -21 - 21 1879 1732 1584 1436 1286 -20 - 22 1134 0982 0828 0674 0518 -19 - 23 0360 0202 0043 982 9721 -16 - 24 9558 9394 9229 9062 8895 -14 - 25 8726 8577 8386 8214 8041 -12 - 27 7867 7691 7515 7337 7158 -8 - 27 6979 6798 6616 6433 8248 -1 - 21 5899 5501 5311 -2 - 30 5120 4928 4735 4541 84346 +1 - 31 <td></td> <td>6336</td> <td>15</td> <td>-12</td> <td>6234</td> <td>6131</td> <td>6027</td> <td>5922</td> <td>5815</td> <td>-10</td> <td>-14</td> <td>5706</td> <td>5597</td> <td>5486</td> <td>5374</td> <td>-107.5</td>		6336	15	-12	6234	6131	6027	5922	5815	-10	-14	5706	5597	5486	5374	-107.5
4059 -19 -18 3932 3803 3674 3543 3411 -20 -19 3278 3143 3007 2870 -2870 -2872 -2882		5261		-15	5146	5030	4913	4795	4675	- 15	-16	4554	4432	4309	4184	-120.2
2732 -20 -19 2593 2453 2311 2168 2024 -21 -21 1879 1732 1584 1436 136 136 1438 1436 1438 1436 1438 1436 1438 1438 1436 1438 1438 1438 1438 1438 1438 1438 1438 1438 1438 1438 1438 1438		4059	- 19	- 18	3932	3803	3674	3543	3411	-20	-19	3278	3143	3007	2870	-132.7
1286 -20 -21 1134 0982 0828 0674 0518 -19 -23 0360 0202 0043 982 9721 -16 -24 9558 9394 9229 9062 8895 -14 -25 8756 8386 8214 8041 -12 -27 7867 7691 7515 7337 7158 -8 -27 6979 6798 6616 6433 6248 -4 -8 6063 5877 5689 5501 5311 -2 -30 5120 4928 4735 4541 4346 +1 -31 4150 3953 3754 3555 +3 -3 3153 2950 2747 2542 2336 +7 -33 1713 1502 +9 -9 +9 1080 0867 0653 0438		2732	-20	- 19	2593	2453	2311	2168	2024	-21	-21	1879	1732	1584	1436	-144.6
9721 -16 -24 9558 9394 9229 9062 8895 -14 -25 8726 8557 8386 8214 8041 -12 -27 7867 7691 7515 7337 7158 -8 -27 6979 6798 6616 6433 6248 -4 -28 6063 5877 5689 5501 5311 -2 -30 5120 4928 4735 4541 4346 +1 -31 4150 3953 3754 3555 +3 -32 3153 2950 2747 2542 2336 +7 -33 2130 1922 1713 1503 1922 +9 -34 1080 0867 0653 0438		1286			1134	0982	0828	0674	0518	-19	-23	0360	0202	0043	9882	-156.5
1 -12 -27 7867 7691 7515 7337 7158 -8 -27 6979 6798 6616 6433 3 -4 -28 6063 5877 5689 5501 5311 -2 -30 5120 4928 4735 4541 4 +1 -31 4150 3953 3754 3555 3355 +3 -32 3153 2950 2747 2542 5 +7 -33 2130 1922 1713 1503 1292 +9 -34 1080 0867 0653 0438	0.99		-16	-24	9558	9394	9229	9062	8895	-14	-25	8726	8557	8386	8214	-168.0
-4 -28 6063 5877 5689 5501 5311 -2 -30 5120 4928 4735 4541 +1 -31 4150 3953 3754 3555 3355 +3 -32 3153 2950 2747 2542 +7 -33 2130 1922 1713 1503 1292 +9 -34 1080 0867 0653 0438		8041	-12	-27	7867	7691	7515	7337	7158	8	-27	6269	8649	6616	6433	-179.3
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		6248	4-	-28	6063	5877	5689	5501	5311	-2	-30	5120	4928	4735	4541	-190.2
+7 -33 2130 1922 1713 1503 1292 +9 -34 1080 0867 0653 0438		4346	+	-31	4150	3953	3754	3555	3355	+3	-32	3153	2950	2747	2542	-201.0
		2336	+7	-33	2130	1922	1713	1503	1292	4	-34	1080	1980	0653	0438	-211.5

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								3	32.	W	A	TE	R	F	- V	-T	D	A'	ΓA		
-221.8	-231.9	-241.8	-251.5	-261.0	-270.4	-279.5	-288.5	-297.4	-306.1	-314.7	-323.1	-331.4	-339.5	-347.6	-355.5	-363.4	-371.0	-378.7	-386.1	-393.6	
8230	5921	3512	1007	8406	5712	2926	0046	7084	4032	0894	7671	4365	8460	7510	3962	0337	6634	2855	9001	5073	
8455	6156	3758	1262	8671	2986	3208	0341	7384	4341	1211	7997	4700	1320	7860	4321	0703	2008	3236	9390	5469	
<u>8</u> 679	6390	4002	1516	8934	6258	3490	0632	7684	4649	1528	$\bar{8}322$	5033	1662	8210	4678	1068	7381	3617	9778	5864	
8903	6624	4245	1769	9116	6530	3771	0922	7983	4956	1844	<u>8</u> 647	5366	2003	8559	5035	1433	7753	3997	0165	6229	
+14 -36	+17 -38	+19 -40	+19 -42	+18 -44	+16 -46	+11 -47	+6 -49	+1 -50	-5 -51	-12 -53	-16 -54	-22 -55	-25 -56	-24 -57	-22 -58	-15 -59	-5 -60	+13 -61	+35 -61	-62	
9125	6856	4487	2021	9458	6801	4051	1211	8281	5263	2159	8970	2698	2343	8907	5391	1797	8125	4376	0552	6653	
9346	7088	4729	2272	9718	7071	4330	1499	8278	5569	2473	$\overline{9}293$	6028	2682	$\bar{9}254$	5746	2160	8495	4754	0938	7046	
9567	7318	4969	2522	9978	7340	4608	1786	8874	5874	2787	9615	6359	3020	<u>9</u> 600	6100	2522	8865	5132	1323	7439	
9286	7547	5208	2771	0237	2092	4886	2072	9169	6177	3099	9936	8899	3358	9766	6454	2883	9234	5508	1707	7830	
0004	7776	5447	3019	0494	7875			9463	6480	3411				0291	6807	3244	9603	5884	2090	8221	
+12 -35				+19 -43	+17 -44		+10 -48	+4 -49	-3 -51	-8 -52	-14 -53	-19 -54	-23 -55	-25 -57	-23 -58	-19 -59			+24 -61		-62
0.998 0221			3266	0751	0.996 8141	5437	2642	0.995 9757		3722	0575	0.994 7344	4030	0635	0.993 7159		0.992 9970		2473	0.991 8612	4675
						_	~~		_		•				~	_	~	_	_		•

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		Lai	ne 93.	-(00	nunue	1)				
IV.	M. Thiesen. ^{200a}	Inc	luded	in I.0	$C.T.^{27}$	2				
$t_1 \rightarrow t_2$	0	1	2	3	4	5	6	7	8	9
40°	0.99 224	186	147	10,	066	024	$\bar{9}8_2$	$\bar{9}4_0$	$\overline{8}9_{6}$	852
50	0.98 807	762	715	669	62_{1}	573	52 ₅	475	425	375
60	0.98 324	272	220	16,	113	05,	005	95₀	894	83 ₈
70	0.97 781	72_{3}	668	60,	548	48,	429	368	307	245
80	0.97 183	121	057	994	93 ₀	86₅	80 ₀	734	$\overline{6}6_8$	$\overline{60}_{1}$
90	0.96 534	467	39,	330	261	19_{2}	122	051	981	90 ₉
100	0.95 838									

Table 93 .- (Continued)

(Cont'd from p. 198)

regarded as a perfectly definite, homogeneous substance, the same the world over. And this, in spite of certain observations indicating the contrary, some of which will be mentioned presently.

Since the discovery of the isotopes all this has changed. We now know that water is not a simple substance, the same everywhere, but is a mixture in which the relative amounts of the several constituents vary with the source and with the manner of purification. Fortunately, this variation is so small in the purified waters commonly used in the study of the properties of water that its effect upon the observed values of those properties is entirely negligible in most cases, thus justifying one in speaking, as in this compilation, of the properties of the ordinary water substance. But in those few cases in which extreme precision of measurement has been attained—in which errors from other sources do not exceed a part in a million or thereabouts—it is necessary to consider whether differences in the composition of different specimens of "pure water" may cause significant differences in the property being studied.

Such a property is the density of water. Over the range 0 to 40 °C values are published to one part in 10 million. But there are as yet no data that enable one to say with certainty whether the density of the "pure water" commonly used in such work is definite to that precision. It probably is not.

About 25 years ago, A. B. Lamb and R. E. Lee ²⁶⁶ reported that the densities of various samples of distilled water, expected to be identical, might differ by as much as 8 parts in 10⁷, even when great care was taken. This was long before the discovery of the isotopes.

If a sample of water contained 1 D_2 to 6500 H_2 —the average ratio ²⁶⁷—then removing the D_2 would decrease its density by about 17 parts in a million. W. H. Christiansen, R. W. Crabtree, and T. H. Laby ²⁶⁸ have reported that the density of rain-water is reduced by 12.7 parts in 10⁶ by

Table 94.—Specific Volume of Compressed Water at a Pressure of 1 Atmosphere

Derived from the densities as given in the corresponding sections of Table 93, where references and comments will be found.

Specific volume = v^* , temperature is $(t_1 + t_2) = t \, {}^{\circ}C$.

In the second column the complete value of v^* is given or indicated; in the following columns, only the last three or four digits, the preceding digits being understood to be those in the left-hand section of the second column, either on or above the line, unless there is a line over the first of the tabulated digits, in which case the immediately preceding digit will differ by one unit from that just specified. For example, Mohler finds $v^* = 1.002715$ at -12° C and 1.003078 at -13° C.

It is interesting to notice that, whereas the specific volume of water at $4 \,^{\circ}$ C is 1.000000 ml/g (=1.000027 cm³/g) when under a pressure of one normal atmosphere, it is 0.999973 ml/g (=1.000000 cm³/g) when the pressure is 1.53 atm (see Table 105).

Unit of $v^* = 1$ ml/g = 1.000027 cm³/g. Temp. = $(t_1 + t_2) = t$ °C.

I. J. F. Mohler. 290

$t_1 \rightarrow t_2$	0	-1	-2	-3	-4 - v*	-5	-6	-7	-8	-9
0	1.000 132	227	327	447	620	825	0 51	282	501	754
- 10	1.002 069	369	715	$\overline{0}78$						

II. C. Despretz.²⁸⁰

III. Revised Chappuis table, 1937. In the subsidiary columns C and T to the right of the v^* values in columns $t_1=0.0$ and $t_1=0.5$ are given the amounts, in units of the last place tabulated, by which each of the corresponding values for $t_1=0.0$ and $t_1=0.5$ in the tables published by Chappuis and by Thiesen, Scheel, and Diesselhorst, respectively, exceeds that here tabulated. Over the intermediate 0.5 °C range these differences may be linearly interpolated. Thus both of those tables may be recovered from this, and so may be the one in the *International Critical Tables*, that being merely the average of the other two. Example: At 17.3 °C this table gives $v^*=1.0012501$; the C value is 9, the T value is 27, and the I.C.T. value is 18 units in the last place greater than that. Whence the C, T, and I.C.T. values are, respectively, 1.0012510, 1.0012528, and 1.0012519, agreeing exactly with those in the tables.

					787	Table 94 (Continued,	ontinuea							
	0.0			0.1	0.2	0.3	4.0	0.5			9.0	0.7	8.0	0.0
		ပ	H						υ	Ę				
1 000	1324	-5	0	1257	1192	1129	1067	1001	-3	7	0949	0893	0839	0786
	0735	-2	ī	9890	0638	0593	0549	0507	-2	-2	0466	0428	0390	0355
	0322	-1	-2	0500	0260	0231	0204	0179	0	-1	0156	0134	0114	0005
	0078	0	0	0063	0020	0038	0028	0019	-	0	0012	000	0003	000
	0000	0	0	0001	0004	8000	0014	0021	-1	+1	0030	0040	0052	9900
	0081	0	+1	8600	0116	0136	0157	0180	0	+2	0204	0230	0258	0287
	0317	0	+3	0349	0382	0417	0454	0492	0	+3	0531	0572	0614	0658
	0703	0	+4	0750	0798	0847	8680	0951	+1	+5	1005	1060	1117	1175
	1235	+1	9+	1296	1358	1422	1487	1554	+1	+1	1622	1692	1763	1835
	1908	+	6+	1983	2060	2138	2217	2297	-1	6+	2379	2462	2547	2632
	2720		+10	2808	2898	2989	3082	3176	-2 -	11	3271	3368	3466	3565
	3665	+2	+13	3767	3869	3974	4080	4187		+14	4296	4405	4516	4628
	4741		+15	4856	4972	2089	5208	5328	+15 +	+15	5448	5571	5694	5819
	5945	+18	+17	6072	6201	6330	6461	6594		+18	6727	6862	2669	7134
	7273	+30	+19	7412	7553	7695	7838	7982		+21	8128	8275	8423	8572
	8722		+22	8874	9076	9180	9335	9491		+23	9649	9807	1966	0128
1.001	0290		+24	0453	0617	0783	0920	1117	+13 +	+26	1286	1456	1628	1800
	1974	+11	+ 56	2148	2324	2501	2679	2858		+28	3038	3220	3402	3586
	3770	+5	+29	3956	4143	4331	4520	4711		+30	4902	5094	5288	5483
	2678	7	+31	5875	6073	6272	6472	6673		+32	6875	7079	7283	7488
	7695	-8 +33	+33	7902	8111	8321	8532	8743	- 10	+34	8956	9170	9385	9601

1818	4138	6558	7106	1694	4406	7212	0111	3101	6180	9349	2604	5946	9372	2883	6476	0152	3908	7744	1660	5654	
1592	3901	6312	8821	1428	4131	6928	9817	2798	5869	9028	2275	2008	9056	2528	6113	9780	3528	7357	1265	5251	
1366	3666	9909	8566	1163	3856	6644	9524	2496	5558	8708	1946	5271	8680	2174	5751	9410	3150	6971	0871	4849	
1142	3431	5822	8311	0899	3582	6361	9232	2194	5247	8389	1618	4934	8336	1821	5390	9040	2772	6585	0477	4448	
- 15 +36	-18 +38	-20 +40	-20 + 42	-19 +44	-17 +45	-12 + 47	-7 +49	-2 + 50	+5 +52	+11 +53	+15 +54	+19 +55	+23 +56	+23 +57	+21 +59	+14 +59	+3 +60	-14 + 62	-36 +63	+63	
$\bar{0}919$	3198	5578	8028	0636	3310	6409	8940	1894	4938	8071	1292	4599	7992	1469	5029	8672	2396	6200	0084	4047	
0697	2962	5335	7805	0374	3038	5798	8650	1595	4630	7754	9960	4264	7649	1117	4669	8304	2020	5816	9692	3647	
0475	2734	5094	7554	$\overline{0}112$	2767	5517	8361	1296	4322	7437	0640	3930	7306	0767	4310	7937	1644	5433	9301	3248	
0255	2503	4853	7304	9852	2498	5238	8072	6660	4016	7122	0316	3598	6965	0417	3952	7570	1270	5051	8911	2850	
$\overline{0}036$	2274	4614	7054	9593	2229	4960	7785	0702	3710	6807	9993	3266	6624	8900	3595	7205	9680	4669	8521	2453	
	-16 + 38			-20 +43	-18 +45	-13 +47	-10 + 52	-5 +51	+2 +51	+8 +53	+14 +54	+17 +55	+22 +56	+24 +57	+22 +58	+19 +59	09+ 6+	-5 +61		-50 +63	
9818	2045	4375	9089	9335	1961	4682	7498	0406	3405	6493	0496	2934	6285	9720	3239	6840	0524	4288	8132	2056	6058
1.001	1.002				1.003			1.004				1.005			1.006		1.007			1,008	
21	22	23	24	25	26	27	28	50	30	31	32	33	34	35	36	37	38	39	40	41	42

Table 94.—(Continued)

$\mathbf{I}\mathbf{V}$	M	Thiesen,	290a	T	CT	272
4 V .	TAT.	THICSCH,				

$t_1 \rightarrow t_2$	0	1	2	3	4	5	6	7	8	9
40	1.00 782	82,	86,	90,	943	98,	Ō2₅	ō7₂	Ī1 ₆	<u>1</u> 6,
50	1.01 207	254	301	34,	39 ₈	448	498	54 ₈	60 ₁	652
60	70 ₅	75_{8}	813	86,	923	97,	$\overline{0}3_{6}$	<u>0</u> 9,	$\bar{1}5_1$	21 ₀
70	1.02 270	33 ₀	39_0	452	513	57 ₆	63,	70 ₈	768	838
80	89,	96₅	$\overline{0}3_2$	0 9,	$\overline{1}6_8$	$\overline{2}3_7$	30₀	376	44,	51 ₈
90	1.03 59 ₀	663	736	710	884	95,	$\overline{0}3_{5}$	$\overline{1}1_1$	$\overline{1}8_{8}$	26₅
100	1.04 343									

(Continued from p. 202)

removing its D_2 , and that in the fractional distillation of tap water the first and the last fractions differed in density by 20.0 parts in 10^6 . Whence they concluded "that, if precise relative determinations of the density of water which had been repeatedly distilled had been made at any time since accurate thermometry has been available, they would have disclosed the fact that natural water is not a simple substance." It seems probable that they intend the reader to understand that the distillation was to be fractional.

J. Mendelejev ²⁶⁹ has reported that the density of the purified water from Lake Baikal increases with the depth from which the sample was drawn, water from 1650 meters being 56 in 10⁷ greater in density than that from the surface. This indicates a gravitational separation of the constituents. (Before distillation, the difference in the densities was about 120 in 10⁷, over twice that after distillation.)

Variability in Water.

In the preceding paragraphs we have considered possible differences between different samples; here we consider possible changes in the same sample, changes arising from other factors than the existing temperature and pressure. The volume of a solution is not, in general, equal to the sum of the volumes of the solute and the solvent. This well-known fact does not concern us now. Some, accepting the idea that water is a mixture of polymers, have advanced the idea that the relative numbers of the several polymers can be disturbed, at least temporarily, but for relatively long periods, by various means, such as antecedent heating, chilling or freezing (see Section 25, Establishment of equilibrium). And it is conceivable that very minute amounts of a soluble impurity may markedly change the polymerization. But no evidence that any of these hypothetical effects are actually of practical significance has come to my attention.

(Go to p. 225)

Table 95.—Specific Volume of Compressed Water: Pressure Exceeding 1 Atmosphere

(For sea-water see Table 108)

The table is divided into the following sections:

- I. Amagat, 0 to 198 °C, 1 to 1000 atm, 13 temperatures, steps of 25 or 50 atm.
- II. Amagat, 0 to 49 °C, 1 to 3000 atm, 10 temperatures, steps of 100 atm.
- III. Bridgman, -20 to +80 °C, 1 to 12 000 atm, 13 temperatures, steps of 500 atm.
- IV. Bridgman, -20 to +100 °C, 1 to 12 000 kg*/cm², 11 temperatures, steps of 500 or 1000 kg*/cm².
- V. Tammann and Jellinghaus, -14 to +15 °C, 1 to 1500 kg*/cm², 26 temperatures, steps of 100 kg*/cm².
- VI. Smith and Keyes, 0 to 360 °C, 1 to 350 atm, every 10°, steps of 25 or 50 atm.
- VII. Tammann and Rühenbeck, 20 to 650 °C, 1 to 2500 kg*/cm², 9 temperatures, steps of 100 kg*/cm².
 - VIII. Adams, 25 °C, 1 to 12 000 bars, steps of 500 or 1000 bars.
 - IX. Trautz and Steyer; reference only.

In every case, values at 1 atm have been retained or inserted so as to facilitate comparisons with the preceding tables.

In the first 5 sections the specific volume has been indicated by the amount $(10^{-4}\Delta)$ by which it falls short of 1 ml/g; in the other three sections the specific volume is given directly. Except in the last section, successive differences have been printed, in distinctive type, between the values from which they have been derived. These differences serve several purposes. They show directly irregularities in the "run" of the values, some of which are disturbingly great; they show at once how the mean temperature coefficient of expansion at constant pressure, and the mean compressibility at constant temperature, each for a tabular step, vary throughout the range covered by the section, and facilitate their evaluation at any point in the table; and they furnish one more means for comparing the results obtained by different observers.

Unit of $v^*=1$ ml/g=1.000027 cm³/g (for this table the distinction between the ml and the cm³ is entirely negligible); of P=1 atm=1.03323 kg*/cm²=1.01325 bars; of p=1 kg*/cm²; of $p_b=1$ bar. Temp. = t °C

I. E. H. Amagat.²⁰¹ His values are expressed in terms of the specific volume at $0 \,^{\circ}$ C and 1 atm; those here given were obtained by multiplying each of his by 1.0001319 so as to reduce them to the same basis as that of Tables 93 and 94; $v^* = 1 - 10^{-4} \Delta$.

²⁹¹ Amagat, E. H., Ann. de Chim. et phys. (6), 29, 68-136, 505-574 (1893).

Table 95.—(Continued)

(Lines continued on p. 209)

t→ P	0		5		10		15 — A-		20		30		40	
1		-1.2			-2.7			00	-17.7	25.7	12.4	22.6	- 77.0	42.5
1	-1.3 12.6	-1.2	-0.1 12.3	+2.6	12.0	6.0	11.9		11.8	45.1	-43.4	33.0	~77.0	74.3
25		0.0	+12.2	2.9	+9.3	6.1			-5.9					
23	12.9	-0.9	12.4	2.9	12.3	0.1	12.0		11.9		23.6		22.5	
50		-0.4	24.6	3.0	21.6	6.4			+6.0	25.8	-19.8	34.7	-54.5	42.2
20		0.7		5.0		0.7				20.0	27.0			
	12.7		12.1		11.8		11.6		11.4					
75	36.9	+0.2	36.7	3.3	33.4	6.6	26.8	9.4	17.4					
	12.5		12.0		11.7		11.4		11.3		22.1		22.3	
100	49.4	+0.7	48.7	3.6	45.1	6.9	38.2	9.5	28.7	26.4	+2.3	34.5	-32.2	42.3
	12.3		11.9		11.6		11.3		11.2					
125	61.7	1.1	60.6	3.9	56.7	7.2	49.5	9.6	39.9					
123	12.2	1.1		3.9		1.2		9.0	11.1		21.7		21.9	
450			11.9		11.5	~ 4	11.3			27.0		24 2	- 10.3	427
150	73.9	1.4	72.5	4.3	68.2	7.4	60.8	9.8	51.0	27.0	24.0	34.3	- 10.5	44.7
	12.2		11.8		11.5		11.2		11.0					
175	86.1	1.8	84.3	4.6	79.7	7.7	72.0	10.0	62.0					
	12.1		11.7		11.4		11.1		10.9		21.5		21.1	
200	98.2	2.2	96.0	4.9	91.1	8.0	83.1	10.2	72.9	27.2	45.5	34.7	+10.8	42.5
	23.8		22.8		22.6		22.0		21.5		21.2		20.9	
250	122.0	3.2	118.8	5.1	113.7	8.6	105.1	10.7	94.4	27.7	66.7	35.0	31.7	42.4
200	23.7	3.2	22.5	3.1	22.3	0.0	21.9	10.7	21.4	<i>D.</i>	20.9	0510	20.5	
300	145.7	4.4	141.3	5.3	136.0	9.0	127.0	11.2	115.8	28.2	87.6	35.4	52.2	42.2
	23.2		22.3		22.0		21.6		21.2		20.6		20.3	
350	168.9	5.3	163.6	5.6	158.0	9.4	148.6	11.6	137.0	28.8	108.2	35.7	72.5	42.3
	22.7		22.2		21.5		21.1		20.8		20.3		20.2	
400	191.6	5.8	185.8	6.3	179.5	9.8	169.7	11.9	157.8	29.3	128.5	35.8	92.7	42.5
	22.5		22.0		21.2		21.0		20.7		20.2		19.6	
450						40.0				20.0		26.4		42.2
450	214.1	6.3	207.8	7.1	200.7	10.0	190.7	12.2	178.5	29.8	148.7	36.4		42.2
	22.1		21.9		21.0		20.5		20.1		19.9		19.9	
500	236.2	6.5	229.7	8.0	221.7	10.5	211.2	12.6	198.6	30.0	168.6	36.4	132.2	42.3
	21.5		210		20.8		20.4		19.9		19.3		19.3	
550	257.7	7.0	250.7	8.2	242.5	10.9	231.6	13.1	218.5	30.6	187.9	36.4	151.5	42.0
	21.3		21.0		20.1		19.8		19.7		19.2		19.2	
600	279.0	7.3	271.7	9.1	262.6	11.2	251.4	13.2	238.2	31.1	207.1	36.4		42.2
200	21.2		20.5	7.2	19.9	11.0	19.6	10.0	19.4	01.1	19.2	00.,	18.9	
650	300.2	8.0	292.2	9.7	282.5	11.5	271.0	13.4	257.6	31.3	226.3	36.7		42.4
050	300.2	0.0	272.2	9.7	202.3	11.5	271.0	13.7	237.0	31.3	220.3	30.7	109.0	72.7
	20.5		20.3		19.6		19.2		19.1		18.7		18.6	
700	320.7	8.2	312.5	10.4	302.1	11.9	290.2	13.5	276.7	31.7	245.0	36 .8	208.2	42.5
	20.4		19.7		19.4		19.0		19.0		18.3		18.4	
750	341.1	8.9	332.2	10.7	321.5	12.3	309.2	13.5	295.7	32.4	263.3	36.7	226.6	42.6
	20.1		19.7		19.2		18.9		18.7		18.3		18.3	
800	361.2		254.0		240.7		200 4		24.4.4	22.0	004 4	26.00	244.0	49.77
800		9.3	351.9	11.2	340.7	12.6	328.1	13.7	314.4	32.8	281.6	36.7	244.9	42.7
050	19.8		19.2		19.0		18.5	40.0	18.3		18.2		17.8	46 -
850	381.0	9.9	371.1	11.4	359.7	13.1	346.6	13.9	332.7	32.9	299.8	37.1	262.7	42.5
	19.3		18.7		18.6		18.4		17.8		18.1		17.5	
900	400.3	10.5	389.8	11.5	378.3	13.3	365.0	14.5	350.5	32.6	317.9	37.7	280.2	42.5
					18.4		17.8		17.7		17.6		17.2	
950					396.7	13.9	382.8	14.6	368.2	32.7	335.5	38.1	297.4	42.7
							17.7		17.5		17.2		17.1	·
1000							400.5	14.8	385.7	33.0	352.7	38.2	314.5	42.3
							200.0		300.1	30.0	554.1	20.0	J . 2.0	

Table 95.—(Continued)

(Lines continued from p. 208)

50		60		70		80 A -		90		100		198
-119 5	49 6	-169 1	56 5	-225 6	63 2	-288 8	68 0	-356 8	74 4	-431 2		
22 8 96 7	49 3	23 1 -146 0	56 2	23 4 -202 2						24.8 -406 4	1130	- 1536
22 2		22 7		23 4						25 0		46
-74 5	48 8	-123 3	55.5	-178 8						-381.4	1109	-1490
21 5		22 0		22 4						24.5		46
-53 0	48 3		55 I	- 156 4	614	-2178				-356 9	1087	-1444
21 3		21 5		22 3		22 9				24 2		46
-317	48 1	-798	543	-134 1	60 8	-1949				-3327	1065	-1398
210		21 0		218		22 4				237		44
-107	48 1	-588	535	-1123	60 2	-1725				- 309 0	1045	-1354
20 7		20 8		21 3		22 0				23 6		44
+100	48 0	-380	530	-910	59 5	- 150 5	65 1	-215 6	698	- 285 4	1025	-1310
20 2		20 4		20 6		217		22 4		23 2		42
30 2	47 8	-176	528	-704	58 4	-1288	64 4	-193 2	69 0	- 262 2	1006	-1268
20 0		20 3		20 4		21 0		22 2		22 8		40
50 2	47 5	+27	527	-500	57 8	-1078	63 2	-171 0	68 4	-239 4	989	-1228
199		20 0		20 2		20 9		217		22 3		39
70 1	47 4	22 7	525	-298	57 1	-869	62 4	-149 3	67 8	-217 1	972	-1189
198		194		198		20 3		212		22 1		37
89 9	47 8	42 1	521	-100	56 6	-666	615	-1281	66 9	- 195 0	957	-1152
19 ó		19 3		197		20 1		20 8		21 5		37
109 5	48 1	61 4	517	+97	56 2	-46 5	608	-107 3	66 2	-173 5	942	-1145
190		19 3		194		20 1		20 3		20 9		36
128 5	47 8	80 7	516	29 1	55 5	-264	60 6	-870	65 6	-152 6	926	-1079
187		19 2		19 1		196		20 0		20 6		35
147 2	47 3	99 9	517	48 2	55 O	-68	60 Z	-670	65 O	-132 0	912	- 1044
18 5		188		188		19 Z		197		20 7		34
165 7	47 0	1187	517	67 0	546	+124	597	-473	640	-1113	899	-1010
18 3		18 4		187		190		193		198		34
184 0	46 9	137 1	514	85 7	<i>54 3</i>	31 4	59 4	-280	63 5	-91 5	884	976
18 2		18 1		18 5		188		190		19.5		33
202 2	47 0	155 2	510	104 2	540	50 2	59 2	-90	63 0	-720	871	943
180		180		18 3		18 2		188		19 5		32
220 2	47 0	173 2	507	122 5	54 1	68 4b	58 6	+98	62 3	-52 5	859	-911
17 5		17 7		18 0		17 9		18 6		190		32
237 7	46 8	190 9	50 4	140 5	54 2	86 3	57 9	28 4	619	-33 5	846	-879
17 0		17 6		17 8		17 9		18 2		187		31
254 7	46 Z	208 5	50 Z	158 3	54 1	104 2	57 6		61 4	-148	833	-848
17 5		17 6		17 8		17 5		17 7		18 5		30
272 2	46 I	226 1	50 0	176 1	54 4	121 7	57 4	64 3	60 6	+37	822	-818

Table 95.—(Continued)

Π.	ы	H. Aı	Amagat.		From the	e same		source a	as the		preceding,	and	treated	in the	same 1	treated in the same manner.	v* ==	1	10-⁴∆.
1.	0.00		2.10		4.35		6.85		10.10		14.25		20.40		29.45		40.45		48.85
. =	-1.3	-1.2	-0.1	1.0-	0.0	+0.6	9.0-	+2.2	-2.8	+4.8	-7.6	+10.9	-18.5	+23.2	-41.7	+37.0	-78.7	+35.4	-114.1
	50.5	,	50.3	,	50.2	,	48.8		47.5		47.3		47.2		45.4		40.4	;	43.8
8	+49.2	-1.0	+20.7	0.0	+20.5	2.0	+48.2	3,5	+44.7	2.0	+39.7	11.0	+28.7	25.0	+3.7	36.0	-32.3	38.0	- 70.3
	49.0		48.0		47.0		47.0		46.5		45.5		45.0		43.5		42.0		43.0
200	98.2	0.0	98.2	+1.0	97.2	2.0	95.2	4.0	91.2	0.0	85.2	11.5	73.7	26.5	47.2	37.5	+6.4	37.0	-27.3
	47.5		46.0		45.5		45.0		44.0		43.0		42.5		42.5		41.0		42.5
300	145.7	+1.5	144.2	1.5	142.7	2.5	140.2	5.0	135.2	7.0	128.2	12.0	116.2	26.5	89.7	39.0	50.7	35.5	+15.2
	45.5		45.0		44.5		43.5		43.0		42.5		41.5		41.5		40.5		41.0
400	191.2	2.0	189.2	2.0	187.2	3.5	183.7	5.5	178.2	7.5	170.7	13.0	157.7	26.5	131.2	40.0	91.2	35.0	56.2
	44.5		43.5		42.5		42.5		42.5		40.5		40.5		39.5		40.0		39.5
200	235.7	3.0	232.7	3.0	229.7	3.5	226.2	5.5	220.7	9.5	211.2	13.0	198.2	27.5	170.7	39.5	131.2	35.5	95.7
	43.0		43.0		45.0		41.5		40.5		40.0		39.0		38.0		38.0		38.5
009	278.7	3.0	275.7	4.0	271.7	4.0	267.7	6.5	261.2	10.0	251.2	14.0	237.2	28.5	208.7	39.5	169.2	35.0	134.2
	41.5		41.5		41.5		40.5		40.0		39.5		38.5		37.5		37.0		37.5
700	320.2	3.0	317.2	4.0	313.2	5.0	308.2	2.0	301.2	10.5	290.7	15.0	275.7	29.5	246.2	40.0	206.2	34.5	171.7
	41.0		41.0		40.5		39.0		38.5		38.5		38.0		36.5		35.5		36.0
800	361,2	3.0	358.2	4.5	353.7	6.5	347.2	7.5	339.7	10.5	329.2	15.5	313.7	31.0	282.7	41.0	241.7	34.0	207.7
	39.5		39.0		38.0		37.0		37.5		37.5		36.0		35.5		34.5		35.0
006	400.7	3.5	397.2	5.5	391.7	7.5	384.2	7.0	377.2	10.5	366.7	17.0	349.7	31.5	318.2	45.0	276.2	33.5	242.7
	38.0		36.5		37.0		36.5		36.5		36.0		35.0		35.0		33.5		34.0
1000	438.7	5.0	433.7	5.0	428.7	8.0	420.7	7.0	413.7	0.11	402.7	18.0	384.7	31.5	353.2	43.5	309.7	33.0	276.7
	35.5		36.0		35.0		36.0		35.0		35.0		34.0		34.0		33.0		33.0
1100	474.2	4.5	469.7	0.0	463.7	7.0	456.7	8.0	448.7	0.11	437.7	19.0	418.7	31.5	387.2	44.5	342.7	33.0	309.7
	35.0		35.0		35.0		35.0		34.5		34.5		34.0		34.0		32.5		32.0
1200	509.2	4.5	504.7	0.0	498.7	2.0	491.7	8.5	483.2	0.11	472.2	19.5	452.7	31.5	421.2	46.0	375.2	33.5	341.7
	34.6		34.1		34.1		34.1		33.5		32.5		32.0		32.5		32.0		31.5
1300	543.8	2.0	538.8	0.0	532.8	2.0	525.8	8.0	516.7	12.0	504.7	20.0	484.7	31.0	453.7	46.5	407.2	34.0	373.2
	33.5		33.0		33.0		33.0		32.1		32.1		31.5		31.5		31.5		31.0
1400	577.3	5.5	571.8	0.0	565.8	2.0	558.8	10.0	548.8	12.0	536.8	20.4	516.2	31.0	485.2	46.5	438.7	34.5	404.2
	32.5		33.0		32.5		31.5		31.5		31.0		31.6		31.0		31.0		30.5

	2	0 703	y	2003	0	600	9	6003	,	0 173	000	0 112	;		;		,	
	£ 5	ن _م ا	3	31.5	•	31.5	70.0	31.0	(.51	30.5	0.02	31.0	57.4	30.0	40.3	31.1	0.00	30.0
•	6	36.3	6.5	629.8	8.0	621.8	10.5	611.3	13.0	598.3	19.5	578.8	32.0	546.8	46.0	500.8	35.9	464.7
		30.5		30.5		31.0		30.5		30.0		29.5		29.0		30.0		29.5
6.5	_	8.999	6.5	660.3	7.5	652.8	11.0	641.8	13.5	628.3	20.0	608.3	32.5	575.8	45.0	350.8	36.4	494.2
		30.0		29.5		29.5		0.62		29.5		29.0		28.5		29.0		28.6
6.5		8.969	7.0	8.689	7.5	682,3	11.5	8.029	13.0	657.8	20.5	637.3	33.0	604.3	44.5	559.8	37.0	522.8
		29.5		29.0		28.5		28.5		29.0		28.5		27.5		28.0		28.0
6.5		726.3	7.5	718.8	8.0	710.8	11.5	699.3	12.5	8.989	21.0	665.8	34.0	631.8	44.0	587.8	37.0	550.8
		29.0		29.0		28.0		28.5		28.0		28.0		27.0		27.5		27.5
6.5		755.3	7.5	747.8	0.0	738.8	0.11	727.8	13.0	714.8	21.0	693.8	35.0	658.8	43.5	615.3	37.0	578.3
		28.5		28.0		28.0		28.0		27.5		27.0		26.0		26.5		597
7.0		783.8	8.0	775.8	0.0	8.992	11.0	755.8	13.5	742.3	21.5	720.8	36.0	684.8	43.0	641.8	37.0	604.8
		27.5		27.0		27.5		27.5		5.92		27.0		26.0		26.0		26.5
7.0		811.3	8.5	802.8	8.5	794.3	0.11	783.3	14.5	768.8	21.0	747.8	37.0	710.8	43.0	8.199	36.5	631.3
		26.5		26.5		27.5		27.5		26.0		26.0		25.5		26.0		26.0
7.0		837.8	8.5	829.3	7.5	871.8	0.11	810.8	16.0	794.8	21.0	773.8	37.5	736.3	42.5	693.8	36.5	657.3
		26.0		26.0		26.0		26.0		25.5		25.0		25.5		25.5		25.5
7.5		863.5	8.5	855.3	7.5	847.8	0.11	836.8	16.5	820.3	21.5	8.864	37.0	761.8	42.5	719.3	36.5	682.8
		26.0		26.0		25.0		25.0		25.0		24.5		25.0		25.0		25.0
7.0		886.8	8.5	881.3	8.5	872.8	0.11	861.8	16.5	845.3	22.0	823.3	36.5	786.8	42.5	744.3	36.5	8.404
		24.5		25.5		24.5		24.5		25.0		24.5		24.5		24.5		24.5
7.5		914.3	7.5	8.906	9.5	897.3	11.0	886.3	16.0	870.3	22.5	847.8	36.5	811.3	42.5	768.8	36.5	732.3
		24.5		24.0		24.0		24.0		24.5		24.0		24.5		24.0		24.0
8.0		938.8	8.0	930.8	9.5	921.3	0.11	910.3	15.5	894.8	23.0	871.8	36.0	835.8	43.0	792.8	36.5	756.3
		24.0		24.0		24.0		23.5		23.5		23.5		24.0		23.5		23.5
8.0		962.8	8.0	954.8	9.5	945.3	11.5	933.8	15.5	918.3	23.0	895.3	35.5	826.8	43.5	816.3	36.5	779.8
		23.0		23.0		23.5		23.5		23.5		23.0		23.0		23.0		23.5
8.0		985.8	8.0	977.8	0.0	8.896	11.5	957.3	15.5	941.8	23.5	918.3	35.5	982.8	43.5	839.3	36.0	803.3
		23.0		24.0		23.5		23.0		23.0		23.0		23.0		22.5		22.5
7.0		1008.8	7.0	1001.8	9.5	992.3	12.0	980.3	15.5	964.8	23.5	941.3	35.5	905.8	44.0	861.8	36.0	825.8

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III. Bridgman.** See p. 214.		*		+16	+791 +7 784 11 144 +7 142	9 926 13	8 1050 16	15	12									

1830 49 1879 1927 48 1975 2021 46 2021 46	80	+73	+1318	1387	65 1452	1514	1572	1626 53	1679	85. 85. 84.	1777	1825	1871	1917
4 4 43			+38	38	98 1452 62	35	36	36	36	35	35	35	35	35
1 73 48 1921 47 1968	02	+72	+1356	1425	39 1490	1551	1608	1662	1715	8 4 4 4 4 4	1812	1860 46	1906	1952
50			+40	30	39	39	39	30	38	38	300	37	37	37
1)22 49 1971														
			+39	30	38 1529	38	38	38	37	37	37	38	38	38
	50	+72	+1435	1503	40 1567	1628	1685	1739	1790	1839	1887	1935	1981	2027
			+40	39	40	9	40	40	40	40	40	40	40	40
			029	7000	7500	8000	8500	0006	9500	10000	10500	11000	11500	12000
	08	-290	47.	+113	167	429 131	560	682	794	900	888	1084	1167	1245
		+63	57	53	50	47	47	45	44	45	42	44	42	39
	02	-277	-17	+166	164	146 426 131	209	727	838	104	1038	1128	1209	1284
		+57	48	52	51 330	20	48	46	45	44	41	38	39	40
	9	-170	201 +31	218	163	526 226	655	773	883	103	93 1079 78	1166	85 1248	76 1324
		+49	51	47	45	43	15	41	40	37	38	39	30	39
	8	-121	203 +82	183 265	43 426 45	569	697	814	923	100	1117	1205	82 1287	76 1363
		£ +	49	4	. 4	43	4	45	45	4	43	41	40	40
9500 10000 10500 11000 12000	1	٠	200	1000	1500	2000	2500	3000	3500	4000	4500	2000	5500	0009

Table 95—(Continued)

III. P. W. Bridgman, (ICT)²⁷⁶ with corrections indicated by the errata published with Vol. 7. Based on Bridgman Proc. Amer. Acad. Arts Sci. 48, 307-362 (1913). As the values he accepted for the specific volumes at 1 atm are somewhat smaller than those in Table 94, all those for a given temperature have been increased by the same amount, so chosen as to make the value at 1 atm the same as that in Table 94. For the temperatures -15 and -20 °C, no value is recorded for 1 atm; values for these temperatures have been increased by 0.0001. His data have been criticized by Tammann,²⁰² partly on the ground that both the direct observations of Amagat and others and the data derived from solutions indicate that, for pressures exceeding 300 kg*/cm², the temperature at which the density is a maximum lies below that at which water and ice are in equilibrium, whereas Bridgman's observations indicate that it lies above that equilibrium temperature even when the pressure is as great as 1500 kg*/cm²; and partly on the shape of the isopiestics at temperatures below 0 °C. (See also Fig. 3 and Section IV.) Bridgman's table in the *Proceedings* (loc. cit.) contains values for every 5 °C, the pressure being expressed in kg*/cm². In the International Critical Tables and in this work, the pressure is expressed in atm. $v^* = 1 - 10^{-4} \Delta$.

Data on pages 212, 213; Fig. 3 on p. 230.

IV. P. W. Bridgman.²⁷⁸ The following values of v^* have been obtained from his table of molecular volumes by dividing by 18.0154. He states: "The results now found do not check in fine detail with those found before; in particular the minimum and maximum of volume as a function of temperature on this isobar at 1500 kg found in 1912 ²⁹²ⁿ and shown in Fig. 40 of the 1912 paper, has not been found this time." In fact, the isobars defined by these values more nearly resemble those found by Tammann than did the earlier ones (see Fig. 3). On comparing these values with those of 1912 (from which those in Section III were derived) systematic differences, often amounting to 2 or 3 units in the next to the last place in Δ (i.e., to 2 or 3 ml/kg), are found. For example, at 50 °C and 4, 5, 6, 7, 8, 9, 10, 11 and 12 kg*/cm² these values exceed the 1912 ones by 1.0, 1.8, 2.2, 2.7, 3.2, 3.3, 3.1, 2.9, and 3.3 ml/kg, respectively.

	Unit of p =	1 kg*/cm ² ;	of $P =$	1 atm	; v* = :	1 - 10	-4 ∆ ml/g	. Т	emp. = t	,C	
$t \rightarrow p$	P	-20	-15		-10	_ A _	-5		0		+20
1	0.9678								- 1	17	- 18
									231		199
500	483.9								+230	49	+181
									194		187
1000	967.8						434	10	424	56	368
							172		167		156
1500	1451.8		630	10	620	14	606	15	591	67	524

²⁰² Tammann, G., and Schwarzkopf, E., Z. anorg. allgem. Chem., 174, 216-224 (1928); Tammann, G., and Jellinghaus, W., Idem. 174, 225-230 (1928).

				Table	95	-(Contin	węd)					
$t \rightarrow p$	P	- 20		- 15		- 10		-5		0	,	+20
•				156		152		148		148		148
2000	1935.7	797	11	786	14	772	18	754	15	739	67	672
		142		134	_ •	131		130		129		129
2500	2419.6	939	19	920	17	903	19	884	16	868	67	801
				121		120		116		117		115
3000	2903.5			1041	18	1023	23	1000	15	985	69	916
				108		106		108		106		100
3500	3387.4			1149	20	1129	21	1108	17	1091	75	1016
						100		98		97		96
4000	3871.4					1229	23	1206	18	1188	76	1112
						175		172		173		179
5000	4839.2					1404	26	1378	17	1361	70	1291
										150		144
6000	5807.0									1511	76	1435
$\stackrel{t\rightarrow}{p}$	P	20		40		50	_ Δ -	60		80		100
		10		70	42	101		471	112	004	151	425
1	0.9768		61	- 79	42	- 121	50	- 171	113	- 284	131	- 435
500	483.9	199	61	199 +120	26	215	F 2	212	101	221 - 63	120	252 183
300	403,9	+ 181 <i>187</i>	01	174	26	+94 162	5 3	+41 173	104	180	120	190
1000	967.8	368	74	294	38	256	42	214	97	+117	110	+7
1000	JU1.0	156	• •	156	•	154		154	,,	159		167
1500	1451.8	524	74	450	39	411	43	368	92		102	174
1000	110110	148	• •	142	•	139		140		142		147
2000	1935.7	672	80	592	42	550	42	508	90	418	97	321
		129		126		127		127		129		134
2500	2419.6	801	83	718	41	677	42	635	88	547	92	455
2000		115	•••	115		116		117	•	119		121
3000	2903.5	916	83	833	40	793	41	752	86	666	90	576
		100	•	105	••	105		106		109		112
3500	3387.4	1016	78	938	40	898	40	858	83	775	87	688
	••••	96		96		96	••	98		99	-	104
4000	3871.4	1112	78	1034	40	994	38	956		874		792
	00/111	179	. •	170		170		170		177		180
5000	4839.2	1291	87	1204	40	1164	38	1126		1051		972
		144		151		152		153		155		157
6000	5807.0	1435	80	1355	39	1316	37	1279		1206		1129
-	0001110		-	130		133		135		135		140
7000	6774.9			1485	36	1449	35	1414		1341		1269
				119		120		122		125		127
8000	7742.7			1604	35	1569	33	1536	70	1466	70	1396
3000				109		110		110		112		114
	0710 6			4740		4.500		4 5 4 5				

202a Bridgman, P. W., Proc. Amer. Acad. Arts Sci., 48, 307-362 (1913).

9000

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1713 34 1679 33 1646 69

1578 *68*

1510

		Ta	ble	95	(Contin	iued)			,		
<i>t</i> → <i>p</i>	P	20	40		50	_ A _	60		80		100
•	•		101		102	_	102		104		105
10000	9678.4	18	314	33	1781	33	1748	66	1682	67	1615
			96		95		95		96		98
11000	10646.2	19	910	34	1876	33	1843	65	1778	65	1713
			84		86		87		88		88
12000	11614.1	19	994	32	1962	32	1930	64	1866	65	1801

V. G. Tammann and W. Jellinghaus.²⁹³ The published values have been multiplied by 1.00013 so as to convert them into ml/g. As these authors assign no value to the specific volume at 1 atm when t is below 0 °C, values taken from Table 94 have been inserted, enclosed in parentheses. The values in this section do not suffice to determine more closely than ± 4 °C the temperature that corresponds to the maximum density along any isopiestic, but they do show the general progressive change in the isopiestics (see Fig. 3). $v^* = 1 - 10^{-4} \Delta \text{ ml/g}$.

t→ p	-8		-7		-6		5 _ Δ		-4		-3	
1	(-15)	-2	(-13)	-3	(-10)	-2	(-8)	-2	(-6)	-2	(-4)	-1
100												
200												
300											145	+1
											48	
400									193	0	193	+2
									43		42	
500							237	1	236	1	235	1
							43		40		44	
600					281	1	280	4	276	-3	279	+2
					43		43		46		42	
700			327	3	324	1	323	1	322	1	321	2
			41		42		33		42		41	
800	371	3	368	2	366	10	356	-8	364	2	362	2
	42		43		42		50		40		40	
900	413	2	411	3	408	2	406	2	404	2	402	9
	38		39		40		40		40		40	
1000	451	1	450	2	448	2	446	2	444	2	442	9

²⁰⁰⁸ Tammann, G., and Jellinghaus, W., Z. anorg. allgem. Chem., 174, 225-230 (1928).

32

Table 95—(Continued)

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<i>t</i> →	-8		-7		-6		5		-4		-3	
Þ							Δ					
	39						33					
1100	490		11				479				10	
	23						28					
1200	513		б				507				8	
	42						39					
1300	555¢		9				546				19	
	34						30					
1400	589•		13				576				21	
	36						32					
1500	6250		17				608				9	
$t \rightarrow p$	-2		-1		0		Δ +1		2		3	
1	(-3)	- 1	(-2)	-1	-1	0	-1	-1	0	0	0	-1
•	(-3)			•		v				•		•
			51		51		52		50		51	_
100			49	-1	50	-1	51	+1	50	-1	51	0
			49		47		46		47		46	
200	98	0	98	+1	97	0	97	0	97	0	97	+1
	46		45		46		46		46		45	
300	144	+1	143	0	143	0	143	0	143	+1	142	1
	47		47		46		45		44		44	
400	191	+1	190	+1	189	+1	188	+1	187	+1	186	1
	43		42		42		43		42		42	
500	234	2	232	1	231	0	231	2	229	1	228	1
	43		43		43		42		43		42	
600	277	2	275	1	274	1	273	1	272	2	270	2
	42		42		41		41		40		39	
700	319	2	317	2	315	1	314	2	312	3	309	1
	41		40		39		38		38		39	
800	360	3	357	3	354	2	352	2	350	2	348	3
	33		38		40		40		41		40	
900	393	-2	395	1	394	2	392	1	391	3	388	2
	40		42		40		39		37		38	•
1000	433	-4	437	3	434	3	431	3	428	2	426	2
				-	35	-				_		
1100					469					12		
					30							
1200					499					9		
-200					28					,		
1300					527					1		
1300					28					1		
1400												
1400					555					1		
					44							

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M-11.	-	(Continu	٠,٠
TRUE	45	(. ontinu	en I

t-	4		5		6		7		8		9	
<i>p</i>	+1	0	+1	-1	+2	-1	+3	+2	+	1 +1	0	+1
	50		50		47		46			7	48	
100	51	0	51	2	49	0	49	1	4	8 0	48	1
	45		45		46		45		4	15	44	
200	96	0	96	1	95	1	94	1	9	3 1	92	2
	45		45		46		46		4	15	44	
300	141	0	141	0	141	1	140	2	13	8 2	136	2
	44		42		43		43		3	8	43	
400	185	2	183	-1	184	1	183	6	17	6 -3	179	2
	42		43		41		42		4	3	41	
500	227	1	226	1	225	0	225	6	21	9 -1	220	3
	41		40		39		37		4	1	38	
600	268	2	266	2	264	2	262	2	26	0 2	258	z
	40		40		41		39		3	9	38	
700	308	2	306	1	305	4	301	2	29	9 3	296	2
	37		37		40		40		3	4	40	
800	345	2	343	-2	345	4	341	8	333	3 –3	336	3
	41		41		38		38		42		36	
900	386	2	384	1	383	4	379	4	375	3	372	2
	38		37		36		38		38		37	
1000	424	3	421	2	419	2	417	4	413	4	409	2
					38							
1100					457							
					33							
1200					490							
					36							
1300					526							
					28							
1400					554							
					13							
1500					567							
$t \rightarrow$	10		11		12			13		14		15
Þ						Δ						
1	-1	0	-1	0	-1	+1		-2	0	-2	+3	-5
	48		48		46			46		45		45
100	47	0	47	2	45	1		44	1	43	3	40
	43		41		42			42		41		43
200	90	2	88	1	87	1		86	2	84	1	83

Table 95—(Continued)

$t\rightarrow$	10		11		12	Δ	13		14		15
<i>p</i> 300	134	1	133	2	131		130	2	128	2	126
	43		42		42		41		41		41
400	177	2	175	2	173	2	171	2	169	2	167
	40		41		39		40		41		40
500	217	1	216	4	212	1	211	1	210	3	207
	39		37		38		37		35		36
600	256	3	253	3	250	2	248	3	245	2	243
	38		39		40		39		40		40
700	294	2	292	2	290	3	287	2	285	2	283
	39		38		38		37		37		37
800	333	3	330	2	328	4	324	2	322	2	320
	37		37		36		36		35		35
900	370	3	367	3	364	4	360	3	357	2	355
	37		37		36		38		38		37
1000	407	3	404	4	400	2	398	3	395	3	392
									34		
1100		28							429		
									37		
1200		24							466		
									29		
1300		31							495		
									37		
1400		22							532		
									22		
1500		13							554		

VI. L. B. Smith and F. G. Keyes.²⁹⁴ These values have been taken from their table, which was computed by means of an empirical equation set up by themselves as a satisfactory representation of their observations. In that table the values of v^* are given to a unit in the sixth place of decimals. But they state that their equation "may be trusted to represent the behavior of liquid water to at least one part in 2000" (p. 294) and that the computed specific volume for 4 °C and one atmosphere differs by "one part in 6900 from the accepted value" (p. 295). In view of these statements it seemed justifiable to give here only 4 places of decimals, corresponding to an accuracy of at least one in 10,000. It will be noticed that the values given for P = 1 atm do not all agree with those in Table 94.

²⁰⁴ Smith, L. B., and Keyes, F. G., *Proc. Amer. Acad. Arts Sci.*, 69, 285-312 (1934) → *Mech. Eng.*, 56, 92-94 (1934). Supersedes Keyes, F. G., and Smith, L. B., *Mech. Eng.*, 53, 132-135 (1931).

Table 95-(Continued)

VI. Smith and Keyes. See p. 219.

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Table 95—(Continued)

VII. G. Tammann and A. Rühenbeck.²⁹⁵ Their values, expressed in terms of the specific volume at 20 °C and 1 atm, have been multiplied by 1.00177 in order to bring them to the same basis as Tables 93 and 94. Both temperature and pressure are carried far beyond the values at the critical point, 374.15 °C, 225.65 kg*/cm², but all the values of the specific volume except one are smaller than the critical volume.

	Un	it of p =	1 kg*/	cm2; of P	= 1 ats	n ; of $v^* =$	1 ml/g.	Temp. =	t °C	
$t \rightarrow p$	P	20		100				300		400
1	0.96	8 1,0018								
		57								
100	96.8	0.9961	382	1.0343	1092	1.1435				
		46		51		85				
200	193.6	0.9915	377	1.0292	1058	1.1350				
		45		51		71				
300	290.3	0.9870	371	1.0241	1038	1.1279				
		43		39		86				
400	387.1	0.9827	<i>375</i>	1.0202	991	1.1192	2442	1.3635		
# 00	1000	44		40		70		158		
500	483.9	0.9783	379	1.0162	961		2354	1.3477	6277	1.9754
		43		27		58		111		339
600	580.7	0.9740	395	1.0135	930	1.1065	2301	1.3366	5049	1.8415
		24		28		46		75		447
700	677.5	0.9716	391	1.0107	912	1.1019	2272	1.3291	4677	1.7968
		43		27		31		77		364
800	774.2	0.9673	407	1.0080	908	1.0988	2226	1.3214	4390	1.7604
		44		28		58		54		371
900	871.0	0.9629	423	1.0052	878	1.0930	2230	1.3160	4073	1.7233
		30		27		57		76		164
1000	967.8	0.9599	426	1.0025	848	1.0873	2211	1.3084	3985	1.7069
		40		8		58		75		152
1100	1064.6	0.9559	458	1.0017	798	1.0815	2194	1.3009	3908	1.6917
		30		27		45		76		204
1200	1161.4	0.9529	461	0.9990	780	1.0770	2163	1.2933	3780	1.6713
		<i>30</i>		16		43		65		218
1300	1258.1	0.9499	475	0.9974	753	1.0727	2141	1.2868	3627	1.6495
		31		28		44		56		217
1400	1354.9	0.9468	478	0.9946	737	1.0683	2129	1.2812	3466	1.6278
		30		16		45		64		168
1500	1451.7	0.9438	492	0.9930	708	1.0638	2110	1.2748	3362	1.6110
		30		17		31		48		164
1600	1548.5	0.9408	<i>505</i>	0.9913	694	1.0607	2093	1.2700	3246	1.5946
		30		15		18		5 2		151

Table 95—(Continued)

					. ,					
t→ p	P	20		100		200 v*		300		400
1700	1645.3	0.9378	520	0.9898	691	1.0589	2059	1.2648	3147	1.5795
		44		16		31		41		138
1800	1742.0	0.9334	548	0.9882	676	1.0558	2049	1.2607	<i>3050</i>	1.5657
		44		16		31		<i>52</i>		86
1900	1838.8	0.9290	576	0.9866	661	1.0527	2028	1.2555	3016	1.5571
		17		16		32		40		125
2000	1935.6	0.9273	577	0.9850	645	1.0495	2020	1.2515	2931	1.5446
		30		27		31		41		72
2100	2032.4	0.9243 40	580	0.9823 6	641	1.0464 <i>32</i>	2010	1.2474 <i>52</i>	2900	1.5374
	0400.0			_			1000		2052	
2200	2129.2	0.9203 20	614	0.9817 <i>16</i>	615	1.0432 <i>31</i>	1990	1,2422 40	2853	1.5275 <i>59</i>
2300	2225.9	0.9183	618	0.9801	600	1.0401	1981	1.2382	2834	1.5216
2300	2223.9	17	010	16	000	31	1901	41	2034	1.3210
2400	2322.7	0.9166	619	0.9785	585	1.0370	1971	1.2341	2764	1.5105
		16		16		18		40		72
2500	2419.5	0.9150	619	0.9769	583	1.0352	1949	1.2301	2732	1.5033
l→ b	P	400		500		550		600		650
900	871.0	1.7233	7476	2.4709						
,,,,	0.1.0	164		1304						
1000	967.8	1.7069	6336	2.3405	4016	2.7421				
		15 2		916		1385				
1100	1064.6	1.6917	5572	2.2489	3547	2.6036				
		204		676		1090				
1200	1161.4	1.6713	5100	2.1813	3133	2,4946	2915	2.7861	3546	3.1407
1200	1050 1	218	4020	498	2060	771 2.4175	2621	1065 2.6796	2620	<i>1991</i> 2.9416
1300	1258.1	1.6495 217	4820	2.1315 452	2860	2.4173 590	2621	2.0190 828	2620	2.9410 1317
1400	1354.9	1.6278	4585	2.0863	2722	2.3538	2383	2,5968	2131	2.8099
1400	1354.9	1.0278	4383	348	6166	572	4363	736	2131	2.8099 860
1500	1451.7	1.6110	4405	2.0515	2498	2.3013	2219	2.5232	2007	2.7239
1000	110111	164	,,,,,	334	- 170	373		655	200.	773
1600	1548.5	1.5946	4235	2.0181	2459	2.2640	1937	2.4577	1889	2.6466
		151		276		423		510		601
1700	1645.3	1.5795	4110	1.9905	2312	2.2217	1850	2.4067	1798	2.5865
		138		241		356		416		628
1800	1742.0	1.5657	4007	1.9664	2197	2.1861	1790	2.3651	1586	2.5237
		86		218		306		415		539
1900	1838.8	1.5571	3875	1.9446	2109	2.1555	1681	2.3236	1462	2.4698
		125		182		304		447		524

²⁹⁶ Tammann, G., and Rühenbeck, A., Ann. d. Physik (5), 13, 63-79 (1932).

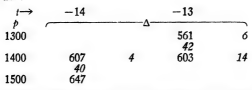
Table 95—(Commed)										
t→ p	P	400		500		550 v*		600		650
2000	1935.6	1.5446 72	3818	1.9264 <i>194</i>	1987	2.1251 209	1538	2.2789 <i>252</i>	1385	2.4174 274
2100	2032.4	1.5374 <i>99</i>	3696	1.9070 <i>160</i>	1972	2.1042 <i>206</i>	1495	2.2537 <i>338</i>	1363	2.3900 <i>357</i>
2200	2129.2	1.5275 <i>59</i>	3635	1.8910 <i>159</i>	1926	2.0836 207	1373	2.2209 285	1334	2.3543 257
2300	2225.9	1.5216 <i>111</i>	3535	1.8751 <i>146</i>	1878	2.0629 189	1295	2.1924 232	1362	2.3286 <i>407</i>
2400	2322.7	1.5105 72	3500	1.8605 <i>136</i>	1835	2.0440 206	1252	2.1692 <i>219</i>	1187	2.2879 <i>306</i>
2500	2419.5	1.5033	3436	1.8469	1765	2.0234	1239	2.1473	1100	2.2573

VIII. L. H. Adams.²⁹⁶ In the conversion of his ratios to specific volumes it has been assumed that the specific volume at 1 atm and 25 °C is 1.0029 ml/g (Table 94).

		Unit of $p_b = 1$	1 bar, of P =	= 1 atm, of v	r = 1 ml/g.	Temp. = 25 °C	D.	
p_b	P	v*	Pb	\boldsymbol{P}	v*	p_b	P	v*
500	493.5	0.9817	4000	3947.7	0.8874	9000	8882.3	0.8166
1000	986.9	0.9635	5000	4934.6	0.8695	9630 ^d	9504.1	0.8098
1500	1480.4	0.9472	6000	5921.5	0.8540	10000	9869.2	0.8059
2000	1973.8	0.9328	7000	6908.5	0.8402	11000	10856.2	0.7964
3000	2960.8	0.9071	8000	7895.4	0.8278	12000	11843.1	0.7876

IX. M. Trautz and H. Stever. 297 These authors have published values for the specific volume of water in the range 50 to 300 atm and 0 °C to near saturation or to 370 °C. M. Jakob 298 regards these values as inferior to those reported by Keyes and Smith in 1931 (see this table, Section VI).

Additional values:



⁴ Equilibrium of water and ice-VI at 25 °C; this pressure "is about 25 bars higher" than the one found by Bridgman.

Specific volume at 750 atm and 100 °C was published as 1.00912, giving $\Delta = 92.5$; here it has been taken as 1.00902.

^b Specific volume at 850 atm and 80 °C was published as 0.99308, giving $\Delta = 67.9$; here it has been taken as 0.99303.

²⁰⁰ Adams, L. H., J. Am. Chem. Soc., 53, 3769-3813 (1931).

²⁰⁷ Trautz, M., and Steyer, H., Forsch. Gebiete Ingenieurw., 2, 45-52 (1931) → Steyer, H., Z. d. Ver. d. Ing., 75, 601 (1930).

²⁰⁸ Jakob, M., Engineering (London), 132, 143-146 (1931).

(Continued from p. 206)

M. Dole and B. Z. Wiener ²⁷⁰ have reported that within their experimental error, which did not exceed one part in 10⁶, the density of a given sample of water is independent of the thermal history of the sample.

But J. B. Peel, P. L. Robinson, and H. C. Smith ²⁷¹ have reported very queer changes in the density of water that has remained for a day or more in contact with carbon or with thoria.

Density of Water.

For the temperature of maximum density, see Section 36. With the exceptions of Tables 108, 109, and 110, referring to natural waters, all the data given below refer to air-free water unless the contrary is stated. The values for the density and the specific volume at a pressure of one atmosphere are exceedingly accurate, but at higher pressures the data obtained by different observers do not always agree satisfactorily.

The values given by V. Stott and P. H. Bigg ²⁷² for the density and the specific volume at one atmosphere and for the range 0 to 40 °C are the means of the corresponding values published by P. Chappuis ²⁷³ and by

(Go to p. 250.)

Table 96.—Isometric Association of the Pressure and Temperature of Compressed Water

Adapted from E. H. Amagat.²⁹⁹ Derived from the same observations as his data in Table 95.

Successive differences have been printed, in distinctive type, between the values from which they have been derived; $v^* = \text{specific volume at } t$ and P, $v_0^* = \text{that at 0 °C}$ and 1 atm. Certain obviously erroneous values in Amagat's tables, arising apparently from errors in transcription, have been changed on the basis of that assumption, so as to smooth the run of the differences. They are marked, and the original values are given in a footnote.

The following isometrics of water at temperatures and pressures close to the critical have been published by C. J. v. Nieuwenburg and Miss H. B. Blumendal ³⁰⁰; unit of $v^* = 1$ cm³/g of P = 1 kg*/cm²:

$t \rightarrow v^*$	350	360	370
1.6	250	334	412
1.8	168	223	286
2.0		189	234

Their values for temperatures and pressures exceeding the critical will be found in Table 36.

²⁰⁰ Amagat, E. H., Ann. de chim. et phys. (6), 29, 68-136, 505-574 (1893). ⁸⁰⁰ v. Nieuwenburg, C. J., and Blumendal, (Miss) H. B., Rec. trav. chim. Pays-Bas, 51, 707-714 (1932).

Table 96—(Continued)

(Lines continued on p. 227)

i→	0		1		2		3		4	
*/5*0						P —				
9778	43.45 50.35	-0.90	42.55 50.90	-0.65	41.90 51.40	-0.28	41.62 51.78	+0.06	41.68 52.11	+
525	93.80 <i>51.3</i>	-0.35	93.45 51.50	-0.15	93.30 51.95	+0.10	93.40 52.32	0.39	93.79 52.6	(
273	145.1 <i>51.9</i>	-0.15	144.95 <i>52.3</i>	+0.30	145.25 52.6	0.47	145.72 53.1	0.68	146.4 53.6	
020	197.0 53.1	+0.3	197.3 53.5	0.6	197.9 53.8	0.9	198.8 <i>54.0</i>	1.2	200.0 54.2	
3766	250.1 53.5	0.7	250.8 54.0	0.9	251.7 54.4	1.1	252.8 54.9	1.4	254.2 55.3	•
513	303,6 54.8	1.2	304.8 55.1	1.3	306.1 55.3	1.6	307.7 55.7	1.8	309.5 56.1	
260	358.4 55.6	1.5	359.9 56.0	1.5	361.4 56.6	2.0	363.4 56.8	2.2	365.6 57.2	2
3007	414.0 56.8	1.9	415.9 57.3	2.1	418.0 57.7	2.2	420.2 58.3	2.6	422.8 58.6	ä
753	470.8 58.1	2.4	473.2 58.4	2.5	475.7 58.7	2.8	478.5 59.0	2.9	481.4 59.5	
499	528.9 59.6	2.7	531.6 59.9	2.8	534.4 60.3	3.1	537.5 60.7	3.4	540.9 60.9	
245	588.5 60.0	3.0	591.5 60.5	3.2	594.7 60.9	3.5	598.2 61.3	3.6	601.8 61.8	
991	648.5 62.2	3.5	652.0 62.5	3.6	655.6 62.9	3.9	659.5 63.1	4.1	663.6 63.5	
736	710.7 62.6	3.8	714.5 62.9	4.0	718.5 63.0	4.1	722.6 63.2	4.5	727.1 63.6	
482	773.3 64.2	4.1	777. 4 64.7	4.1	781.5 65.4	4.3	785.8 66.1	4.9	790.7 66.6	
227	837.5 65.7	4.6	842.1 66.1	4.8	846.9 66.5	5.0	851.9 67.2	5.4	857.3 67.7	
972	930.2	5.0	908.2	5.2	9134	5.7	919.1	5.9	925.0	
→	0		5	10		15	2	0	30	
/v*0 200					P					
50										
00										
50										
25									36.5 21.9	7
15								3.2 55.2	58.4	7
05						5.2).8 i.0 <i>56.8</i>	<i>22.4</i> 80.8	7
						10.2	10	-	11.6	

Table 96—(Continued)

(Lines continued from p 226)

5		6			7	P	8			9		10
42 00	+0 60	42	60 -	+0 90	43 50	+1 25	44 7	5 +1	75	46 50	+2 15	48 65
52 50		52	85		53 20		53 4	5		536		536
94 50	0 95	95	45	1 25	96 70	1 50	98 2	0 1	9	100 1	21	102 2
529		53	2		53 6		54 0	1		54 4		548
147 4 54 1	1 3	148 <i>54</i>		16	150 3 54 7	19	152 2 55 1		3	154 5 55 3	25	157 0 55 5
201 5 54 5	16	203 54		19	205 0 55 3	23	207 3 55 7	2 :	5 :	209 8 56 1	27	212 5 56 5
256 0 55 6	20	258 56	0	23	260 3 56 4	27	263 0 56 6	2 5	, ;	265 9 56 8	3 1	269 0 57 0
311 6 56 4	24	314 56		27	316 7 57 0	29	319 6 57 2	3 1	! :	322 7 57 7	3 3	326 0 57 9
368 0	27	370		30	373 7	3 1	376 8	3 0	5 3	880 4	35	383 9
57 5		58			58 3		58 9			59 2		599
425 5 59 0	3 2	428 59		3 3	432 0 59 5	37	435 7 59 7	3 9	,	439 6 <i>60 1</i>	42	443 8 60 5
484 5	34	487		36	491 5	39	495 4	43	3 4	199 7	46	504 3
60 0		60			610		61 5			618		62 0
544 5 61 2	39	548 61		4 1	552 5 62 0	4 4	556 9 62 2	4 0	5 5	61 5 62 5	48	566 3 62 8
605 7	43	610	0	4 5	614 5	46	619 1	45	,	524 0	5 1	629 1
62 3	, ,	62			62 9		63 6	•		640	-	64 3
668 0 63 8	46	672 (64 .		48	677 4 64 6	5 3	682 7 64 7	5 3	3 (688 0 65 1	5 4	693 4 65 6
731 8	49	736	7	5 3	742 0	5 4	747 4	5 7		753 1	59	759 0
64 1		64 :	5		65 0		65 6			66 1		66 6
795 9	5 3	801 2	2	58	807 0	60	813 0	6 2		319 2	64	825 6
67 0		67 5	5		67 7		67 8			680		68 4
862 9	58	868		60	874 7	61	880 8	6 4		887 2	68	894 0
68 3		68 8	8		69 2		697			70 1		70 3
931 2	63	937 5	5	6 4	943 9	66	950 5	68	9	57 3	7 0	964 3
40		50		60		70 P	F	80		90		100
						51 5	134 5	186 0	146 0	332		485 0 118 0
				38	5 122 0	109 0 160 5	137 5	112 0 298 0	147 0	113 (445 (603 0
				111		114 5	10, 5	117 5		120		122 5
		39 5	110 5			275 0	140 5	415 5	149 5	565		725 5
		1140		118	0	122 0		1130		114	,	118 0
57 0	96 5	153 5	1145	268	0 129 0	397 0	131 5	528 5	151 0	679 5	1640	843 5 ^a
56 5		58 5		61		615		720		75 0		75 5
113 5	98 5	212 0	117 5	329	5 129 0	458 5	142 0	600 5	154 0	754	5 164 5	919 0
22 5	-	24 0		24		25 5		26 0		26 5		27 0
136 0	100 0		117 5			484 0	142 5	626 5	154 5	781		946 0
23 5		24 5		24		25 5		260		26 (27 0
159 5	101 0		117 5			509 5	143 0	652 5	154 5	807 (973 0
11 6		11 5		12		12 5		12 5		13 5		13 5
171 10	100 0		118 5		5ª <i>131 5</i>	522 0	143 0	665 0	155 5	820 5		986 5
	200 7			370				3000		020		,50 5

Table 96—(Continued)
(Lines continued on p 229)

			-	(Lin	es cont	inueu c	ш		,	20		30	
$t \rightarrow v^*/v^*_0$	0		5		10		. P	15		20		30	
	95				10 1			10 6		10 8		11 3	
0 9995	10 5	-21	8 4	5 4	13 8	12 2		26 0	19 3	45 3	58 4	103 7	793
	19 2		196		20 4			20 5		216		22 8	
85	29 7	-17	28 0	62	34 2	12 3		46 5	20 4	66 9	59 6	126 5	<i>80 5</i>
	19 3		20 0		20 3			21 6		22 1		23 0	
75	49 0	-10	48 0	65	54 5	13 6		68 1	20 9	89 0	60 5	149 5	81 5
	50 O		52 2		530			549		56 O		58 O	
50	99 0	+12	100 2	7 5	107 5	<i>15 5</i>		123 0	<i>22 0</i>		62 5	207 5	<i>84 0</i>
	51 O		52 5		54 5			56 1		57 5		60 0	
25	150 0	27	152 7	93	162 0	17 1		179 1	23 4		65 0	267 5	85 5
0 9900	<i>51 0</i> 201 0	47	<i>53 0</i> 205 7	11 1	54 8 216 8	187		56 4 235 5	24 5	57 5	67 0	<i>59 5</i> 327 0	88 0
0 9900	105 4	47	108 5	11 1	112 2	10 /		233 3 115 0	44 3	118 0	07 0	123 0	00 V
0 9850	306 4	78	314 2	148	329 0	21 5		350 5	27 5	378 0	720	450 0	920
0 7000	109 6		1136	2,0	1160			1190		122 0		128 0	,,,
800	416 0	118	427 8	17 2	445 0	24 5		469 5	30 5	500 0	78 0	578 0	96 0
	1130		117 2		121 0			125 0		127 0		132 0	
750	529 0	160	545 0	210	566 0	28 5		594 5ª	32 5	627 0	83 0	710 0	100 0
	118 0		121 0		125 5			127 5		131 0		137 0	
700	647 0	190	666 0ª	25 5	691 5	<i>30 5</i>		722 0	<i>36 0</i>	758 0	89 0	847 0	107 0
	121 5		125 5		130 0			134 5		137 0			
650	768 5	23 0	791 5	<i>30 0</i>	821 5	35 0		856 5ª	38 5	895 0			
	127 0		132 0		133 0			1 3 8 5					
0 9600	895 5	28 0	923 5	31 0	954 5	40 5		995 0					
t→ v*/v ₀ *	0.00	0	10 1	0	20	40 P		2	9 45		40 45		48 85
1 000	1	0			3		4		88	86	174	86	260
	200				22	25		2	32		243		248
0 990	201	15	216	43			51	3	20	97	417	91	508
	217		233		24			2	52		263		266
0 980	418	31	449	51	50	00 7	72	5	72	108	680	94	774
	230		245		20	50		2	73		283		293
0.970	648	46	694	66	70	60 8	85	8	345	118	963	104	1067
	247	-	264			80			89		306		315
0 960	895	63	958	82	104		04		34	135	1269	113	1382
0.00	275		289			04	•		16		327		334
0 950	1170		1247	97	134		96		50	146	1596	120	1716
0 700	295		312	71		24	, 0		30	1 70	342	200	360
	493		314		34			_	50		JTE		500

[°] Published values of the pressure were as follows 5°, 0 9700, P=2660; 15°, 0 9750, P=5985, 15°, 0 9650, P=8505, 40°, 1 0000, P=1715; 60°, 1 0000, P=3955; 60°, 0 9995, P=4050, 100°, 1 0050, P=8535, 0 00°, 0 915, P=2335.

Table 96-(Continued)

				(Line	s conti	nued fi	om p.	228.)				
40		50		60	·····	70 P	······································	80		90		100
11.9		12.5		12.5		13.0		13.5		13.0		13.5
183.0	101.5	284.5	118.5	403.0¢	132.0	535.0	143.5	678.5	155.0	833.5	166.5	1000.0
24.0		25.0		24.5		25.0		26.5		26.5		
207.0	102.5	309.5	118.0	427.5	132.5	560.0	145.0	705.0	155.0	860.0		
24.0		24.0		25.5		26.0		26.0		27.0		
231.0	102.5	333.5	119.5	453.0	133.0	586.0	145.0	731.0	156.0	887.0		
60.5		63.0		64.0		65.5		67.0		69.0		
291.5	105.0	396.5	120.5	517.0	134.5	651.5	146.5	798.0	158.0	956.0		
61.5		63.5		65.0		66.0		68.0				
353.0	107.0	460.0	122.0	582.0	135.5	717.5	148.5	866.0				
62.0		62.5		65.0		67.0		68.5				
415.0	107.5	522.5	124.5	647.0	137.5	784.5	150.0	934.5				
127.0		131.5		135.0		139.0						
542.0	112.0	654.0	128.0	782.0	141.5	923.5						
132.0		136.0		140.0								
674.0	116.0	790.0	132.0	922.0								
136.0		148.0										
810.0	128.0	938.0										
144.0												
954 0												

$t \rightarrow v^*/v_u^*$	0.00		10.10		20.40	P	29.45		40.45		48.85
0.940	1465	94	1599	109	1668	112	1780	158	1938	138	2076
	158		165		171		184		189		191
0.935	1623	101	1724	115	1839	125	1964	163	2127	140	2267
	162		172		179		188		193		197
0.930	1785	111	1896	122	2018	134	2152	168	2320	144	2464
	169		178		189		197		206		2 05
0.925	1954	120	2074	133	2207	142	2349	177	2526	143	2669
	176		183		193		191		199		213
0.920	2130	127	2257	143	2400	140	2540	185	2725	157	2882
	195		204		205		216		228		
0.915	23254	136	2461	144	2605	<i>151</i>	2756	197	2953		
	183		192		209		213				
0.910	2508	145	2653	161	2814	155	2969				

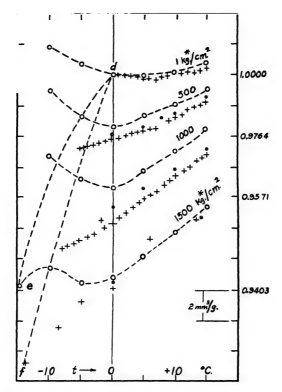


FIGURE 3. Isopiestic Variation of the Specific Volume of Water with the Temperature.

(See p. 214.)

[Adapted from G. Tammann and W. Jellinghaus, Z. anorg. allgem. Chem, 174, 225-230 (1928).] Ordinates represent changes in the specific volume, 1 division - 2 mm³/g; abscissas represent the temperatures, in °C.

For convenience, the several isopiestics have been relatively displaced, crowded together; to the right is given for each isopiestic the specific volume corresponding to the ordinate division next below the lowest point of the Bridgman isopiestic. The dots indicate the observations of E. H. Amagat [Ann. chim. phys. (6), 29, 68-136, 505-574 (1893)]; the circles, those of P. W. Bridgman [Proc. Am. Acad. Arts Sci., 47, 439-558 (1912); 48, 307-362 (1913)]; and the crosses, those of Tammann and Jellinghaus (loc. cit.). The lines de and df connect the several melting points as determined by Bridgman and by Tammann and Jellinghaus, respectively. The pressure corresponding to each isopiestic is indicated.

Table 97.—Isopiestic Thermal Expansion of Water

Adapted from a table computed by L. B. Smith and F. G. Keyes ²⁰⁴ by means of an empirical equation representing their observations. They give the values to five significant figures, but in view of what they say about the limitations of their equation (see Table 95, Section VI), three significant figures seem to be sufficient for this compilation. Similarly, it has not seemed necessary to give here their values for P = 1, 25, 75, 125, and 175 atm; those for P = 1, extending only to t = 100 °C, are, to the precision

Table 97—(Continued)

of this table, identical with those for $P_{\rm sat}$, and the others can be obtained without significant error by interpolation between those here given. Adjacent to each t, is given the corresponding value of the pressure $(P_{\rm sat})$ of the saturated vapor. Each value of $(dv^*/dt)_p$ given under $P_{\rm sat}$ in the body of the table is the limiting value approached along the isopiestic corresponding to $P_{\rm sat}$ as t approaches that corresponding to saturation, *i.e.*, the associated t. The exponent n is given in the third column, and each value is to be used until another is given.

Examples: At 140 °C the saturation pressure is 3.57 atm; along that isopiestic $(dv^*/dt)_p$ at 140 °C is $10.48 \times 10^{-4} = 0.001048$. At 150 °C and P = 50 atm, $(dv^*/dt)_p = 1.10 \times 10^{-3} = 0.00110$.

	U ₁	nit of	P and of	$P_{\text{sat}} = 1$ at	m; of (dv*	$(dt)_p = 1$	nl/g·°C.	Temp. = 1	°C.	
$P \rightarrow t$	Past	n	Pant	50	100	150 - 10 ⁿ (dv*/	200	250	300	350
0 10 20 30 40	0.006 0.012 0.023 0.042 0.073	5	-3.09 +9.39 2.00 2.95 3.80	2.94 3.78	2.94 3.76	2.93 3.74	2.92 3.72	2.91 3.70	2.90 3.68	2.89 3.66
50 60 70 80 90	0.122 0.196 0.308 0.467 0.692		4.57 5.28 5.96 6.61 7.24	4.54 5.24 5.91 6.54 7.16	4.51 5.20 5.85 6.48 7.08	4.48 5.16 5.80 6.41 7.00	4.45 5.12 5.75 6.35 6.92	4.42 5.08 5.70 6.29 6.85	4.39 5.04 5.65 6.23 6.78	4.36 5.00 5.60 6.17 6.71
100 110 120 130 140	1 1.41 1.96 2.67 3.57		7.86 8.49 9.13 9.79 10.48	7.77 8.38 9.01 9.64 10.31	7.67 8.26 8.88 9.49 10.12	7.58 8.15 8.76 9.34 9.95	7.49 8.05 8.65 9.20 9.78	7.40 7.95 8.54 9.06 9.62	7.32 7.85 8.43 8.93 9.46	7.24 7.75 8.33 8.80 9.32
150 160 170 180 190	4.70 6.10 7.82 9.90 12.4	3	1.12 1.20 1.28 1.37 1.48	1.10 1.18 1.26 1.35 1.44	1.08 1.15 1.23 1.31 1.40	1.06 1.13 1.20 1.28 1.37	1.04 1.11 1.18 1.25 1.33	1.02 1.08 1.15 1.22 1.30	1.01 1.06 1.13 1.20 1.27	0.99 1.05 1.11 1.17 1.24
200 210 220 230 240 250	15.3 18.8 22.9 27.6 33.0 39.2		1.59 1.71 1.86 2.02 2.21 2.44	1.55 1.68 1.82 1.98 2.18 2.41	1.51 1.62 1.75 1.90 2.07 2.28	1.46 1.57 1.69 1.82 1.98 2.17	1.42 1.52 1.63 1.76 1.90 2.07	1.38 1.48 1.58 1.70 1.83 1.98	1.35 1.44 1.53 1.64 1.76 1.90	1.32 1.40 1.49 1.59 1.71 1.84
260 270 280 290 300	46.3 54.3 63.3 73.5 84.8		2.71	2.69	2.53 2.83 3.22 3.72 4.43	2.38 2.65 2.98 3.39 3.93	2.26 2.50 2.78 3.13 3.57	2.16 2.36 2.61 2.91 3.28	2.06 2.25 2.47 2.73 3.05	1.98 2.16 2.35 2.58 2.85
310 320 330 340 350	97.4 111 127 144 163				5.46	4.68 5.76 7.63° 10.99°	4.15 4.94 6.08° 7.89° 11.68°	3.74 4.36 5.18 6.47 8.28	3.43 3.91 4.53 5.53 6.72	3.17 3.55 4.03 4.84 5.76

[&]quot;For P = 175, $10^{\circ}(dv^*/dt)_p = 6.68$ at 330 °C, 9.30 at 340 °C, and 14.39 at 350 °C.

Table 98.—Mean Isopiestic Coefficient of Thermal Expansion of Water

For slopes of isopicstics see Table 97. Values for other pressures and ranges in temperature may be readily obtained from the values given in Table 95.

Unit of P = 1 atm = 1.0332 kg*/cm²; of p = 1 kg⁴/cm². Temp. = t °C

I. Pressure = 1 atm. Coefficients have been computed from the data in Table 94. They essentially agree with the earlier observations by G. T. Gerlach,³⁰¹ as quoted in the fifth edition of Landolt-Börnstein's Tabellen.

For α_1 the temperature is expressed in terms of the numbers x; the first line contains the values for each degree from 0 to 9; the second, for each 10° from 0 to 90; the third, for each 10° from 5 to 95. Examples: The mean coefficient between 40 and 41 °C is 38.92×10^{-5} ; that between 45 and 46 °C is 43×10^{-5} .

 $\alpha_1 = 10^5 (v_{t+1} - v_t) / v_t$; $\alpha_{10} = 10^5 (v_{t+10} - v_t) / 10 v_t$.

II. Derived from the same observations as the specific volumes given in Table 95.

A = Amagat; taken from his more detailed table ²⁰¹ which differs slightly from and supersedes his earlier table.³⁰² The latter is still frequently quoted in certain important compilations.

B = Bridgman; computed by the compiler from B's data in Table 95(III).

SK = Smith and Keyes: computed by the compiler from their table of specific volumes, with retention of more significant figures than are given in Table 95.

		Uı	nit of $P =$	1 atm; α, =	= 105 (v2 -	$-v_1)/(t_2-t_1)$	v_1		
Interval→		0° to 10	0	1	10° to 2		30°	to 40°	
Obsciver→	A	В	sk	A	В	SK	A	В	SK
\boldsymbol{P}					$-\alpha_m$				
1	1.4^{a}	2	3.4	14.9	15	14.8	33.4	35	32.4
100	4.3°		3.9	16.5		15.2	34.5		33.5
200	7.2		4.3	18.3		15.3	35.0		33.5
300	9.8		4.8	20.5		15.5	35.7		33.2
500	14.9	12		23.6	27		37.0	37	
700	19.2			26.2			37.7		
1000	25.9	27		29.4	29		39.6	38	
2000	36.4	40		35.6	38		42.3	42	
4000		41			42			44	
6000		3 6			42			47	

⁸⁰¹ Gerlach, G. T., "Salzlösungen," Freiberg, 1859.

³⁰² Amagat, E. H., Compt. rend., 105, 1120-1122 (1887).

140	16 30	Commi	neu j	
		60° to 70	O°	
SK	A	В	SK	A
41.6	rr /	a	rr o	(1

Interval→	4	0° to 5	0°		60° to 7	Oo		70° to 8	00
Observer $\rightarrow P$	A	В	SK	A	В	SK	A	В	SK
1	42.2	43	41.6	55.6	α <u></u>	55.3	61.8	62	61.5
100	42.2		41.3	54.8		54.6			60.6
200	42.6		40.9	53.9		53.9	60.0		59.8
300	42.3		40.6	52.8		53.4	59.0		58.9
500	42.9	50		52.3	48		56.6	57	
700	43.4			52.3			55.4		
1000	43.7	45		51.2	53			54	
2000		46			53			49	
4000		49			49			46	
6000		46			46			45	

III. Comparison of Bridgman's values of 1912 (B') and of 1935 (B"). The B' values have been computed from the specific volumes given in his table of 1912, and the B" ones from those given in Table 95, Section IV. The 1912 table is the basis of the International Critical Table values, which are given in Table 95, Section III.

$$\alpha_m = 10^5 (v_2 - v_1) / (t_2 - t_1) v_1$$

Unit of $p=1 \text{ kg*/cm}^2$; of P=1 atm

Int	erval->		- 20°		-40°		-50°		-60°	60°-		80°-100°
	Set→ n	B'	В"	B'	B"	B'	B"	B'	B"	B'	B"	В"
Þ	P						$-\alpha_m$					
0		8	8	30	30	42	42	49	49	59	56	74
500	483.9	19	24	33	31	44	26	49	53	55	52	60
1000	967.8	27	29	36	38	44	39	49	43	53	50	56
2000	1935.6	36	36	41	43	45	45	47	44	50	47	51
4000	3871.4	41	43	43	44	45	45	46	42	48	46	45
6000	5807	38	45	46	47	45	45	46	43	45	42	44
8000	7743			51		47	42	46	39	44	41	41
10000	9 678					48	40	46	40	45	40	40
12000	11614					49	40	48	40	45	40	40

IV. W. Watson.³⁰³ L. Bouchet ³⁰⁴ thinks that these values of α are vitiated by a systematic error. The pressures are all higher than the critical ($p_{crit} = 225.65 \text{ kg*/cm}^2$), and most of the temperatures exceed t_{crit} (374.15 °C). At the higher temperatures the thermal expansion is several times as great as that of the ideal gas. These coefficients are with reference to the volume (v_0) at room temperature and the indicated pressure. $\alpha' = 10^5 (v_2 - v_1)/(t_2 - t_1)v_0$. For the ideal gas $\alpha' = 341$ if room temperature = 20 °C.

Unit of $p=1 \text{ kg*/cm}^2$; of P=1 atm.

$\begin{array}{c} t_1 \rightarrow \\ t_2 \rightarrow \\ p \end{array}$	P	100 200	200 300	300 400	400 500	500 600	600 700	700 800	800 900
400 700 1000	387.1 677.5 967.8	99 91 84	600 380 280	3280 880 520	4300 1760 1000	1880 1060	1880 1060	1060	1060

^a The specific volume passes through a minimum in the interval here covered.

²⁰⁶ Watson, W., Proc. Roy. Soc. Edinburgh, 31, 456-477 (1911). Briefer report in Ber. Sachs. Ges. Wiss, Leipzig (Math.-Phys.), 63, 264-268 (1911). 804 Bouchet, L., Compt. rend., 178, 554-556 (1924).

Table 99.—Interpolation Formulas for the Thermal Expansion of Water

Of the numerous interpolation formulas that have been proposed for the thermal expansion of water, those more frequently quoted are here assembled. In Table 100 the corresponding values defined by them are compared with one another and with the data of Tables 94 and 260. It will be seen that some of the formulas are entirely unsatisfactory; they should not appear in future hand-books or similar compilations.

For temperatures not exceeding 100 °C, the pressure is assumed to be 1 atm and invariable; for higher temperatures, it is the pressure of the saturated vapor, the water being continuously saturated; v_0 = actual volume at 0 °C, v_t = volume of the same mass at t °C. To each formula is assigned a key symbol serving to suggest its proposer and to identify the associated data in Table 100.

The development of a formula for liquids has been discussed by E. Salzwedel, ³⁰⁵ W. Jazyna, ³⁰⁶ W. Herz, ³⁰⁷ V. Fischer, ³⁰⁸ O. Tumlirz, ³⁰⁹ and others.

I. Formulas of the type $10^6(v_t - v_0)/v_0 = a + bt + ct^2 + dt^3 + et^4$; temp. = t °C.

II. Other formulas. Temp. = t °C.

Keye Formula Range, °C

T-C
$$1-\rho=\frac{v_t-1}{v_t}=\frac{(t-3.9863)^2}{508929.2}\cdot\frac{t+288.9414}{t+68.12963}$$
. 0 to 40

Th' $1-\rho=\frac{v_t-1}{v_t}=\frac{(t-3.98)^2}{503570}\cdot\frac{t+283}{t+67.26}$ 0 to 40

Th" $1-\rho=\frac{v_t-1}{v_t}=\frac{(t-3.98)^2}{466700}\cdot\frac{t+273}{t+67}\cdot\frac{350-t}{365-t}$ 17 to 100

Web $10^6\left(\frac{v_t-v_t}{v_4}\right)=8.2004(4-t)+5.44402(4-t)^2+0.26698(4-t)^3-10$ to +4

ZT $10^6\left(\frac{v_t-v_t}{v_0}\right)=51700+845(t-110)+3.5(t-110)^2$ 110 to 140

Table 99—(Continued)

* Key and references:

Ch', Ch", Ch" Chappuis, P., Trav. et Mém. Bur. Int. Poids et Mes., 13, D.1-D.40 (1907).

Hirn, G. A., Ann. de chim. et phys. (4), 10, 32-92 (1867). La

Landesen, G., Schr. Naturf. Ges. Univ. Jurjeff (Dorpat), No. 14 (1904); based on his observations published in No. 11 (1902).

Panebianco, H., Riv. di. min. e crist. ital., 38, 3-11 (1909); reviewed in Z. Kryst., 50, 496-497 (1912). Pa

M. L., Ann. de chim. et phys. (3), 15, 325-408 (1845); see Frankenheim, M. L., Ann. d. Physik (Poga.), 86, 451-464 (1852). Scheel, K., Idem (Wied.), 47, 440-465 (1892). Pi

Sch

Tilton's improved formulation of the observations of Chappuis; see Tilton, L. W., and Taylor, J. K., J. Res. Nat. Bur. Stand., 18, 205-214 (RF971) (1937).

Thiesen, M., Scheel, K., and Diesselhorst, II., Wiss. Abh. Physik. Techn. Reichsanstalt, 3, 1-70 (1900); Ann. d. Physik (Wied.), 60, 340-349 (1897).

Thiesen, M., Wiss. Abh. Physik. Techn. Reichsanstalt, 4, 1-32 (1903). T-C

Th'

Th" We

Weidner, Ann. d. Physik (Pogg.), 129, 300-308 (1866). Zepernick, K., and Tammann, G., Z. physik. Chem., 16, 659-670 (1895). ZT

^b Both the formulas, Pa and We, are based solely upon the data reported by Weidner; their difference, actually insignificant, results from certain arithmetical errors in Weidner's computation. An examination of Weidner's data shows that they do not justify the retention of more than three significant figures in each of the coefficients of the equation, and probably two would be enough.

In the Landolt-Börnstein Tabellen, the We formula is given incorrectly, the coefficients of the formula in Section II of this table being given as applicable to the formula of Section I. The same error in the 1905 edition of those Tabellen was pointed out by H. Panebianco in 1909, in the paper in which the Pa formula was derived. That formula is now included in the Tabellen as an independent one, but the error in the We formula is perpetuated. Furthermore, the reference given in the Tabellen for the Pa formula is Riv. di min. c crist. ital., 38, 3 (1912); the year should have been given as 1909, 1912 corresponding to vol. 41, in which there is nothing relating to the Pa formula.

Actually published in the equivalent form shown in the second section of the table.

Table 100.—Comparison of Interpolation Formulas for the Thermal Expansion of Water

The several formulas, with their key symbols, are given in Table 99. For $t \ge 100$ °C the pressure is 1 atm; for t > 100 °C the pressure is that of the saturated vapor. In the second column are given the values of $10^6(v_t-v_0)/v_0$ as determined from the data in Tables 94 and 260, the doubtful digits being overscored; v_0 is the volume at 0 °C. By adding to any of the values in column two the appropriate value of δ , the corresponding value of $10^6(v_t - v_0)/v_0$ as defined by the interpolation formula is obtained. Example: The value of $10^6(v_t - v_0)/v_0$ as defined by the Pi formula is 2583 - 210 = +2373 at $12 \,^{\circ}$ C, and +195 + 3.9 = +198.9at -2 °C.

Since the values in column Obs for the range 0 to 40 °C have been derived by the T-C formula, the δ for that formula is zero throughout this range. Parentheses enclose values of δ that lie beyond the range of the formula. It is obvious that formulas Pi, We = Pa, La, Hi, and ZT are quite unsatisfactory.

³⁰⁵ Salzwedel, E., Ann. d. Physik (5), 5, 853-886 (1930).

³⁰⁶ Jazyna, W., Z. Physik, 58, 429-435, 436-439 (1929).

⁸⁰⁷ Herz, W., Z. Elektroch., 32, 460-462 (1926).

⁸⁰⁸ Fischer, V., Ann. d. Physik (4), 71, 591-602 (1923).

²⁰⁰ Tumlirz, O., Sitz.-ber. Akad. Wiss. Wien (Abt. IIa), 130, 93-133 (1921).

Value of 10°(Value of 100(v1-v2)/v2 as defined by formula is Obs+3. Temp1°C	fined by for	rmula is Obs	+6. Temp.	J. 1=								
Key	$\frac{\mathrm{Obs}^{b}}{10^{4}}$	F	We=Pa	T-C	Ch'	Ch"	Ch",	Th'	Th"	Sch	La	Hi	ZZ
	+2583	- 210											
- 10	1937	-252	- 160										
	1369	-216	- 163										
9 -	919	-172	- 163										
- 4	488	- 50	- 74										
- 2	+ 195	+ 3.9	- 29	(-21.9)									
+ 3	-100.2		+ 3.2	0	+0.4			-0.1		+ 5.2			
4	-132.4		- 5.1	0	+0.4			+0.0		+ 7.2			
9	-100.7			0	+0.5			+0.3		+ 7.1			
∞	- 8.9			0	+0.6			+0.6		0.9 +			
10	+ 139.6			0	+0.3	(-0.3)		+1.0		+ 4.4			
10.5	185.2			0	+0.1	+0.3		+1.1		+ 4.0			
11	234.1			0		+1.1		+1.3		+ 3.7			
11.5	286.3			0		+1.4		+1.4		+ 3.2			
12	341.7			0		+1.8		+1.5		+ 2.8			
12.5	400.3			0		+2.0		+1.6		+ 2.5			
13	462.0			0		+2.3	+2.3	+1.8		+ 2.2			
15	739.7			0			+2.4	+2.2	+3.3	+ 1.1			
20	1636.9			0			-0.3	+3.3	+5.1	+ 19			

										+240	+161	+160	+279	(+426)			
									- 150	- 202	- 266	-233	- 209	- 200	- 200	-619 -2017	
(+ 8.5)	+ 1.3	- 1.9	- 3.1	- 2.6	- 14.4	-34.0	-37.2										
+ 4.2	+10.9	(+21.7)															
+6.5	+7.0	+6.4	+4.2	+1.3	4.4	-8.3	-9.2	-4.4	-3.8								
+4.3	+5.2	+5.7	+6.1														
-1.5	+0.7	+2.9	-2.0														
0	0	0	0	(+5.9)													
2800.7	4207.5	5838.8	7679.9	11940	16920	22570	28860	35770	43300	51400	60200	00969	00161	90500	102000	1274 <u>00</u> 1564 <u>00</u>	
22	30	35	4	20	8	20	8	8	100	110	120	130	140	150	160	180	

*These so-called observed values have been derived from the data in Tables 94 and 260; values for t < 0 are from Mohler's data; for t between 0 and 40 °C from those defined by Tilton's formula (T-C), for t between 40 and 100 °C from Thiesen's data, all given in Table 94; and for t > 100 °C from the data in the International Skeleton Steam Table of 1934 (Table 260). • For the significance of the key symbols, except Obs (= observed), see Table 99.

Table 101.—Thermal Slopes of the Isometrics of Water

(For mean isometric thermal coefficient of pressure, see Table 102)

Bridgman's data (B) have been read from his published graphs [P. W. Bridgman ^{292a, Figs. 9, 10}]; the others have been adapted from G. Tammann and A. Rühenbeck.²⁹⁵

 $v_{1,20}$ and $v_{1,0}$ = specific volumes at 1 atm and at 20 °C and 0 °C, respectively; v^* = actual specific volume; pressure = 1 atm + p. $v_{1,20}$ = 1.00177 and $v_{1,0}$ = 1.00013 ml/g.

	2,0		,	0								
			Unit	of $p=1$	kg*/cm	2 = 0.96	78 atm.	Temp.	=t°C			
$t \rightarrow$	0	20	40	60	80		$t\rightarrow$	0	20	40	60	80
Þ			- (dp/dt)v	(B) ——			Þ			$(dp/dt)_v(B)$) ——	
0	-0.5	+4.5	9.1	12.5	15.0		10000			41.0	39.8	36.0
1000	+ 6.0	9.2		15.2	15.8		11000			43.0	41.6	37.5
2000	11.4	13.5	15.2	17.4	17.4		12000			44.1	42.3	37.6
3000	15.7	16.8	18.0	19.5	19.2		$t\rightarrow$	0	20	40	60	80
4000	18.8	19.7	20.6	21.8	21.5		$v^{*/}v_{1,0}$			$(dp/dt)_v(B)$		
5000	20.3	22.0	23.4	24.3	23.6		1.00	-1	+4.1	9.4	13.5	15.2
6000	17.8	25.4	27.0	27.0	26.0		0.95	+7.2	11.0	14.0	17.1	18.0
7000		32.8	30.8	30.2	28.4		0.90	16.1	18.0	19.6	22.0	23.0
8000		38.5		33.2	30.8		0.87_{5}	19.8	21.2	23.8	25.6	26.5
9000			38.0	36.7	33.5		0.85	18.5	27.0	30.0	31.0	31.7
				Та	ımmanı	and	Ruhenbo	ck				
v*/1'1,2		2.1			2.13			2 1	8		2.25	
1	$(d_{I}$	$b/dt)_{u}$	Þ	(dp/	$dt)_{v}$	Þ	(dt	v/dt)v	Þ	(dp/d	$t)_{v}$	Þ
525		4.2	1692	13	.6	1640	13	2.3	1508	10.7		1368
575	1	1.8	2348	12	.0	2278	10	0.7	2028	9.6		1876
625		7.3	281.2	7	.4	2762		8.7	2565	8.3		2323
					•							

Table 102.-Mean Isometric Thermal Coefficient of Pressure of Water

(For slopes of isometrics at 0 to 80 and 500 to 600 °C, see Table 101)

Adapted from E. II. Amagat,²⁹⁹ with correction of certain computational errors and with adjustment to accord with Table 96.

 $\gamma = \frac{1}{P_1} \left(\frac{P_2 - P_1}{t_2 - t_1} \right)_v; \ v = \text{volume of a certain mass of water at } P \text{ and } t,$ $v_0 = \text{that of the same mass at 1 atm and 0 °C.}$

		Unit o	$f_{\gamma}=1 \ ($	°C)-1.	remperat	ures are t_1	°C and	t₂°C		
$\begin{array}{c} t_1 \rightarrow \\ t_2 \rightarrow \\ v/v_0 \end{array}$	10	10 20	20 30	30 40	40 50	50 60 10 ⁴ γ	60 70	70 80	80 90	90 100
1.0200 1.0150 1.0100						280 _o	317 ₀ 834	261 ₀ 857 511	785 493 360	461 355 284
1.0050 1.0025				211 _o	169 ₀ 868	747 554	481 392	331 310	286 256	241 218
1.0015 1.0005 1.000	-27₀	833₀	1725 ₀ 237 ₀ 168 ₀	133 ₀ 974 852	735 633 590	498 451 435	369 348 337	294 281 274	247 237 234	211 203 202

Table	10	2(C	ontinued))
			_		

$\begin{array}{c} t_1 \rightarrow \\ t_2 \rightarrow \end{array}$	0 10	10 20	20 30	30 40	40 50	50 60	60 70		70 80	80 90	90 100
v /vo ,					1	104γ					
0.9995 0.9985	314 152	228 ₀ 956	129 ₀ 891	765 637	555 493	417 381	32 31			229 220	200
0.9975 0.9950 0.9925 0.9900	112 86 80 79	633 349 250 199	680 431 321 258	545 406 320 269	443 360 303 259	358 304 265 238	29- 26- 23- 21-	0 2 3 2		213 198	
0.9850	74	149	190	204	207	196	18	1			
0.9800 0.9750 0.9700 0.9650 0.9600	70 70 69 69 66	124 108 98 89	156 132 117	166 141 126	172 158	167					
$\begin{array}{c} t_1 \rightarrow \\ t_2 \rightarrow \\ v/v_0 \end{array}$	0.00	10.10 20.40	20.40 29.45 - 10 ⁴ γ -	29.45 40.45	40.45 48.85	$\begin{array}{c} t_1 \rightarrow \\ t_2 \rightarrow \\ v/v_0 \end{array}$	0.00	10.10 20.40	20.40 29.45 — 1047	29.45 40.45	40.45 48.85
1.000 0.990 0.980 0.970 0.960 0.950 0.940	75 74 70 69 65 64	194 110 92 83 76 68	1760 258 160 124 100 87 74	889 276 172 127 108 92 81	588 259 165 129 106 90 85	0.935 0.930 0.925 0.920 0.915 0.910	62 62 61 59 58 57	65 62 62 62 57 59	75 73 71 65 64 61	75 71 69 66 65	79 74 67 69

Table 103.—Isothermal Compression of Water

[See also Tables 104 (dilated water), 105 (mean between 1 and P atm.), and 106 (mean between P_1 and P_2 atm.)]

E. Brander ³¹⁰ has found that Bridgman's 1912 values (see Table 95) up to 6000 kg*/cm² can be represented by the empirical expression: $\log_{10}(1 + Ap) = B(v_1 - v_p)/v_p$ where v_1 and v_2 are the volumes of the same mass of water at the temperature considered and at the pressure of 1 and p kg*/cm², respectively, and A and B depend on the temperature only, taking the following values:

1	0	20	40	60	80	•C
500A	0.136	0.111	0.105	0.102	0.106	cm²/kg*
B	2.368	2.1668	2.128	2.084	2.101	dimensionless

C. Grassi,³¹¹ working within the range 1 to 10 atm, found the isothermal compressibility $[\beta = -(1/v)(dv/dp)_t]$ for a given temperature to be independent of the pressure; it varied with the temperature, the following values being reported:

t 0 1.5 4.1 10.8 13.4 18.0 18.0 25.0 34.5 43.0 53.0 °C 10°8 50.3 51.5 49.9 48.0 47.7 46.3 46.0 45.6 45.3 44.2 44.1 per atm

The following values have been selected from a table computed by

⁸¹⁰ Brander, E., Soc. Scien. Fennica, Comm. Phys. Math., 7, No. 7 (1934).

⁸¹¹ Grassi, C., Ann. de chim. et phys. (3), 31, 437-478 (1851) \rightarrow J. de Pharm. et Chim. (3), 19, 442-444 (1851).

Table 103—(Continued)

L. B. Smith and F. G. Keyes ²⁹⁴ by means of an empirical equation representing their observations, and have been rounded off to three significant figures (see remarks at head of Section VI of Table 95) theirs being given to five. They also give values for P=1, 25, 75, 125, and 175 atm. Those for P=1 atm, t > 100 °C are, within the precision of this table, identical with those for $P_{\rm sat}$; those for the other pressures can be satisfactorily obtained by interpolation, except as noted.

Adjacent to each value of t is given the corresponding value of the pressure (P_{sat}) of the saturated vapor. Each value of $(dv^*/dp)_t$ given under P_{sat} in the body of the table is the limit approached along the isotherm as P approaches P_{sat} ; v^* is the specific volume.

	Unit of	P and of Pa	at = 1 atm	; of (dv*/	$(dp)_i = 10^{-6}$	ml/g·atm	. Temp.	=t°C	
$P \rightarrow t$	P_{nat}	Past	50	100	150 (dv	200 */dp): ——	250	300	350
0	0.006	46.1	45.0	44.1	43.1	42.2	41.3	40.5	39.6
10	0.000	45.5	44.5	43.6	42.7	41.8	40.9	40.1	39.3
20	0.023	45.2	44.3	43.4	42.5	41.6	40.8	40 0	39.2
30	0.042	45.2	44.3	43.4	42.6	41.7	40.9	40.1	39.3
40	0.073	45.5	44.6	43.7	42.8	42.0	41.2	40.4	39.6
50	0.122	46.0	45.1	44.2	43.3	42.5	41.7	40.9	40.1
60	0.196	46.8	45.8	44.9	44.0	43.2	42.4	41.6	40.8
70	0.308	47.8	46.8	45.9	45.0	44.1	43.3	42.4	41.7
80	0.467	49.0	48.0	47.1	46.1	45.2	44.4	43.5	42.7
90	0.692	50.6	49.5	48.5	47.6	46.6	45.7	44.8	44.0
100	1	52.4	51.3	50.2	49.2	48.2	47.3	46.3	45.4
110	1.41	54.6	53.4	52.3	49.2 51.2	50.1	49.1	48.1	47.2
120	1.96	57.2	55.9	54.7	53.5	52.4	51.3	50.2	49.2
130	2.67	60.1	58.8	57.5	56.2	54.9	53.7	52.6	51.4
140	3.57	63.8	62.2	60.7	59.3	57.9	56.6	55.3	54.1
150	4.70	67.9	66.1	64.4	62.8	61.3	59.8	58.4	57.0
160	6.10	72.8	70.8	68.9	67.1	65.4	63.7	62.1	60.6
170	7.82	78.5	76.2	74.0	72.0	70.0	68.1	66.2	64.5
180	9.90	87.3	82.6	80.0	77.6	75.3	73.1	71.0	69.0
190	12.4	93.2	90.0	87.0	84.2	81.5	78.9	76.5	74.1
200	15.3	102.8	99.0	95.3	91.9	88.7	85.7	82.8	80.0
210	18.8	114.2	109.6	105.2	101.0	97.1	93.4	90.0	86.7
220	22.9	128.0	122.5	116.9	111.8	107.0	102.5	98.3	94.4
230	27.6	145	138	131	125	119	113	108	103
240	33.0	166	158	148	140	133	126	120	114
250	39.2	192	183	170	159	150	141	133	126
260	46.3	226	217	197	183	170	159	149	139
270	54.3	270		232	212	195	180	168	156
280	63.3			278 °	250	227	208	191	176
29 0	73.5			342 °	300	268	242	220	201
300	84.8			437	371 °	324	287	257	232
310	97.4			623	473 "	401 *	348	306	274
320	111				634 °	514 °	433	374	329
330	127				919	693 •	556	468	406
340	144				1653	992 •	746	603	512
350	163					1618 °	1054	795	651
360	184					3816	1693	1094	831

°Value of (dv^*/dp) , for P=75 is 296 at 280°, 390 at 290°; for P=125, 400 at 300°, 522 at 310°, 730 at 320°; for P=175, 434 at 310°, 567 at 320°, 784 at 330°, 1125 at 340°, and 2318 at 350°.

Table 104.—Mean Isothermal Compressibility of Dilated Water (Adapted from J. Meyer ³¹²)

There is no discontinuity in the value of the compressibility (β_m) as P changes from positive (= pressure) to negative (= tension).

Values of P for which no value of β_m has been determined have been omitted from the table.

 $\beta_m \equiv (v_1 - v_2)/v_1(P_2 - P_1)$; here, $P_1 = 0$, $v_1 = v$ olume at zero pressure and the indicated temperature, $v_2 = v$ olume of the same mass at the same temperature and P_2 ; the water always completely fills the vessel at P_2 and the indicated temperature, v_2 being the volume of the vessel under those conditions.

Uı		atm = 1.01	325 megad	ynes/cm²; o	$f \beta_m = 10^{-6}$	per atm.	Temp. $= t$ °C	
!→	1	2	3	4	5	6	7	8
9	40.2			ρ,	m			46.2 46.7
10 11 12	48.3	48.4	48.2		47.7	47.1	46.7	
13			48.2	47.7	47.8			
19 20				47.6	47.2	47.0 47.0	46.9	
21				17.0	47.4	-17.0		45.4
24 26				47.8	47.6	47.1		47.1
t→ P	9	10	11	12 β	13	14	15	16
	,			F-1	46.3			
2 4 5 6 7				46.7		46.6		46.5
6			46.4		46.6			
8		46.3			40.0		46.8	
8 10		1010		46.5		46.5		
11 12			46.5			40.3		
15	46.0	46.6	467					
18 20	46.9	46.8	46.7					
23	46.9							
i→ P	17	18	19	20	21 m ———	22	23	24
	,				т	45.7		45.3
3					46.8	46.5		
2 3 4 5 6								46.1
0					47.3 47.1		46.2	
7 8					46.8		70.2	
9 10				47.2 47.3			46.2	46.4
12				47.2		46.5		46.5

⁸¹⁹ Meyer, J., Abh. d. Deutsch, Bunsen Ges., 3 No. 1, whole number 6 (1911).

Table 104—(Continued)											
t→ P	17	18	19	20	β _m ———	22	23	24			
-13 -14 -15			47.1		ρ _m	46.5 46.7	46.7 46.8				
-16 -17		47.4	47.2		46.8 47.0	40.7	46.8 47.1	46.2			
-18 -19		47.4		47.0	46.8 46.9			46.3			
-20 -21	47.4	47.5		17.0	10.5	46.9		46.3 46.4			
-22	47.6					47.2	46.3	40.4			
-23 -24						47.3	46.4				
-26 -27					47.4 47.5		10,1				
-29				47.2	47.5						
$_{P}^{t\rightarrow}$	25	26	27	β	29	30	31				
+ 7 + 6 + 4 + 3 + 2	46.1 45.8 46.4			46.0 45.6 45.8	m.			·			
+ 2 - 2		45.4	46.0	46.6		•	45.4				
$-\frac{4}{5}$	46.1	46.5	45.8				45.1				
- 6 - 7		46.0	45.8			45.7	45.1				
- <i>7</i> - 9	46.2 46.1	46.3				45.1 45.2					
$-10 \\ -11$	10.1	46.2				45.3					
$-12 \\ -14$	46.0				45.0	45.7					
-14 -16	46.1				45.8 45.9						
$-18 \\ -20$				45.8	46.0						
$-21 \\ -23$				46.0 45.9							
-26			46.1	43.9							
-29 -30		46.1	46.2								

Table 105.—Mean Isothermal Compressibility of Water Between Pressures 1 and P

(Between P_1 and P_2 , Table 106; adiabatic compressibility, Table 107; natural waters, Table 110).

Values for other temperatures and pressures may be readily obtained from the specific volumes as given in Table 95.

Table 105-(Continued)

These mean compressibilities $[\beta_m = (v_1 - v_p)/(P-1)v_1]$ cannot be satisfactorily represented by an expression of the form $\beta_m = a - bP$, which has been proposed by K. Drucker³¹³ and by A. L. T. Moesveld.³¹⁴ The former gives for 10^6a and 10^6b , respectively, the values 47.0 and 0.0115 at 25 °C, and 46.3 and 0.015 at 35 °C; the latter gives 44.5 and 0.00492 at 25 °C. The unit of P is 1 atm in each case. W. Jazyna ³¹⁵ has considered the calculation of the compressibility from other data.

 $\beta_m = (v_1 - v_p)/(P-1)v_1$ or $(v_0 - v_p)/Pv_0$, where v_0 , v_1 , and v_p are the volumes of the same mass of water at the same temperature but under the pressures 0, 1, and P, respectively. The values of v_1 have been taken from Table 94, unless others have been specified in the article cited. As t varies, β_m passes through a minimum at about 40 or 50 °C for Amagat's and most of Bridgman's observations, but near 30 °C for the values given by Smith and Keyes and for some of Bridgman's more recent data.

Unit of $P=1$ atm, of $P_b=1$ bar,	of $b = 1 \text{ kg*/cm}^2$: of $\beta_m = 10^{-8}$	per unit of pressure.	Temp. = 1 °C.

$\stackrel{t\rightarrow}{P}$	0	10	20	30	40 βm —	50	60	80	100
1 <10	51.5 50.3 50.3	48.4 47.8 48.3	46.4 45.9 46.0	45.3 44.8 45.3	44.8 44.3 44.5	44.7 44.4 44.1	45.0 44.8	46.7 46.1	48.2
25	52.5 45.7	50.0 45.3	49.1 44.9	44.8	44.9	45.2	45.7	47.4	49.9
50	52.0 45.6 50.0	49.6 45.0	48.3 44.7	48.0 44.6	45.5 44.7 43.2	45.9 45.0	46.3 45.5 43.5	47.1 44.8	48.7 49.7
100	51.2 51.0 45.0	48.3 48.0 44.5	46.9 47.5 44.2	46.0 45.7 44.1	44.9 46.5 44.2	44.9 43.7 44.6	45.0 45.0	46.6	49.1
200	50.0 50.0 44.1	47.1 47.2 43.6	45.4 46.2 43.3	44.5 44.5 43.2	43.8 40.1 43.4	43.6 43.1 43.7	44.1 44.2	45.8 45.8	47.4 48.2
300	49.2 49.2 43.2	46.4 46.1 42.7	44.6 45.0 42.4	43.6 43.8 42.4	42.9 43.0 42.5	42.8 42.8 42.9	43.1 43.3	44.9 44.9	46.8 47.2
500	47.6 47.5 46.6	45.0 44.8 44.6	43.3 43.4 42.3	42.3 42.4 41.8	41.6 41.7 41.5	41.4 41.6 40.1	41.6 39.5	43.3 42.0	45.4
1000	44.0 43.4	41.7 41.0	40.3 40.3 39.6	39.5 39.4 38.7	38.9 38.6 38.4	38.7 38.7 38.2	38.9 38.2	39.9 39.2	41.7
2000	38.2 38.0	36.5 36.2	35.6 35.1	34.9 34.6	34.4 34.2	34.2 34.1	34.2	34.4	35.0
3000	33.9 33.6	32.7 32.4	31.9 31.7	31.5 31.1	31.1 31.0	31.0 30.8	30.9	31.5	

⁸¹⁸ Drucker, K., Z. physik. Chem., 52, 640-704 (661) (1905).

⁸¹⁴ Moesveld, A. L. T., Idem, 105, 450-454 (1923).

⁸¹⁵ Jazyna, W., Z. Physik, 58, 858-860 (1929).

		10	
Table	105	(Con	imued)

$t \rightarrow p$	-10	0	10	20	β _m	40	50	60	80	Ref.
500	44.5	45.8 46.4	43.0	41.6 39.7	40.6	40.4 39.4	40.0 40.4	40.0 41.8	41.2 43.2	B B'
1000	41.8	42.2 42.6	39.9	38.5 38.6	37.6	37.3 37.0	37.1 37.2	37.1 37.8	38.0 39.0	B B'
1500	40.0 42.5	3 9.3 3 9.5	37.5	36.3 36.0	35.5	35.2 35.0	35.1 35.0	35.1 35.7	36.0 36.3	B B'
2000	38.6 39.4	37.0 37.0	35.4	34.4 34.4	33.7	33.4 33.3	33.2 33.1	33.4 33.4	34.1 34.2	B B'
3000	34.4 34.6	32.8 32.9	31.7	30.9 31.1	31.1	30.2 30.2	30.1 30.1	30.2 30.2	31.1 30.8	B B'
5000	28.3 28.4	27.4 27.2	26.7	26.2 26.1	26.1	25.8 25.4	25.7 25.4	25.8 25.5	26.2 26.0	B B'
7000			23.3	23.0	23.0	22.5 22.2	22.5 22.2	22.5 22.2	22.9 22.6	B B'
10000					19.3	19.1 18.8	19.1 18.8	19.1 18.8	19.3 19.1	B B'
12000						17.4 17.1	17.4 17.2	17.4 17.2	17.6 17.4	В В'

t = 25 °C. Unit of $P_b = 1$ bar = 106 dynes/cm². 296

								10000	
β_m	42.5	39.3	37.0	35.0	31.5	26.6	23.2	19.6	17.9

a References and remarks:

- A E. H. Amagat. Values derived from Table 95, Section I.
- A' E. H. Amagat. Values derived from Table 95, Section II.
- B P. W. Bridgman. Values in the first section of the table (pressure in atm) derived from Table 95, Section III; those in the second section (pressure in kg*/cm²) derived from his table, soca from which the values in Table 95, Section III, were derived.
- B' P. W. Bridgman.²⁷⁸ Values derived directly from the table there given.
- G C. Grassi. Ent Between 1 and 10 atm he could detect no variation of β_m with the pressure. Values here given were obtained by interpolating between his values of β , which may be found in Table 103.
- J R. S. Jessup. 816 Merely incidental determinations made for the purpose of testing his apparatus.
- R C. R. Randall.⁸¹⁷ Values he derived from his determinations of the adiabatic compressibility (β_a) by means of the formula $\beta = \beta_a + T (dv/dt)^2/vc_p$; even at 86 °C the last term is less than 10 per cent of β_a .
- SK L. B. Smith and F. G. Keyes. 204 Values derived directly from their table of specific volumes.
- T D. Tyrer. SIS Values he deduced from his determinations of the adiabatic compressibility. (See R, above.)

⁸¹⁶ Jessup, R. S., Bur. Stand. J. Res., 5, 985-1039 (RP244) (1930).

⁸¹⁷ Randall, C. R., Bur. Stand. J. Res., 8, 79-99 (RP402) (1932).

²¹⁸ Tyrer, D., J. Chem. Soc. (London), 103, 1675-1688 (1913).

Table 106.—Mean Isothermal Compressibility of Water Between Pressures P_1 and P_2

(For $P_1 = 1$ or 0, see Table 105; for adiabatic compressibility, see Table 107; for natural waters, see Table 110.)

Values for other temperatures and pressures than those here given may be readily obtained from the specific volumes as given in Table 95 and from the more extended original tables from which these have been taken or computed.

A few values for 0 and 30 °C and for irregularly distributed pressures up to 95 atm have been published by Earl of Berkeley, E. G. J. Hartley, and C. V. Burton.⁸¹⁹ They do not differ significantly from those found by others, and are not given in this table.

 $\beta_m = (v_1 - v_2)/(P_2 - P_1)v_1$, where v_1 and v_2 are the volumes of the same mass of water at the common temperature indicated and under the pressures P_1 and P_2 , respectively.

			Unit o	f P = 1 att	m ; of $\beta_m =$	10-6 per	atm.			
$\stackrel{t\rightarrow}{P_1}$	0	10	20	30	β _m	50	60	80	100	
					$P_2 = P_1 + 2$	25				
1	52.5 45.7	50.0 45.3	49.1 44.9	44.8	44.9	45.2	45.7	47.4	49.9	
25	51.6 45.3	49.2 44.8	47.6 44.5	44.4	44.5	44.9	45.4	47.0	48.0	
75	50.2 44.5	47.0 44.0	45.3 43.6	43.6	43.7	44.0	44.6	46.1	48.6	
25	49.1 43.6	46.3 43.1	44 6 42.9	428	42.9	43.3	43.8	45.3	47.7	
175	48.8 42.8	46.0 42.3	43.8 42.1	42.0	42.2	42.5	43.0	44.6	46.9	
				P_{2}	$= P_1 + 10$	0				
0	51.1 44.6	48.3 44.1	46.8 43.8 45.8	46.0 43.7	44.9 43.8	44.9 44.1	45.5 44.6		47.8 48.7	
100	49.2 43.3	46.1 42.8	44.2 42.6 44.8	43.6 42.6	42.9 42.7	42.5 43.0	42.7 43.5	45.1	46.8 47.4	
200	48.0 41.7	45.3 41.3	43.4 41.1 42.4	42.4 41.1	41.4 41.2	41.6 41.6	41.5 42.0	43.6 43.5	45.9 45.7	
400	45.5	43.0	41.5 39.9	40.6	40.4	39.9	39.4	40.8	43.4	
500 800 900	42.9 40.6	40.5 38.9	39.4 37.3 36.5	38.7 37.4 36.0	38.2 36.2 35.3	37.7 36.2 35.3	38.3 36.3 36.0	38.7 36.3 35.7	40.7 38.2 37.1	
				I	$P_2 = P_1 + 5$	00				
1	47.5 46.6 46.5	44.7 44.6 43.6	43.4 42.3	42.4 41.8	41.7 41.5	41.6 40.1	39.5	42.0		

⁸¹⁹ Earl of Berkeley, Hartley, E. G. J., and Burton, C. V., Phil. Trans. (A), 218, 295-349 (1919).

			1	able 1	l0 6 ((Continu	ed)			
$t \rightarrow P_1$	0	10	20	30		0 5		60	80	100
2 1						+500				
500	41.6 41.0 41.6	39.5 38.1 38.8	38.0 37.7	35.5	36. 36.	2 36	.6	37.5	37.1	
1000	35.8 36.0 34.5	34.8 34.8	33.8 33.9		33. 33.		.5 .1	33.3	33.8	
1500	32.4 32.6	31.3 31.2	30.9 30.4	30.1 30.5	30. 30.			30.2	30.6	
2500	26.1 26.1	25.9 26.0	25.7 25.7	25.8 25.4				25.2	25.8	
	20.7	21.1 17.0	21.0	20.0	20		0	20.6		
8000 10000 11500			13.9	11.6	11.	8 11.	.8	13.0 11.7 11.4	11.6	
em per	atures abov	re 100 °C								
$t \rightarrow P_1$	P ₂			40		1 - β _m —			200	220
25	50	53.1 47.6		8.0 1.5	64.8	7	4.0	8	36.4	104.4 83.3
$\stackrel{t\rightarrow}{P_1}$	P_2	240	20	50	280	300 β		320	340	360
	350	97.8	117	7.		184.				584
P_1	190 13.0 52.5	10.8	19.0	23	2	230 27.7 55.6	33 2	2	29.6	46.1
β_m	104	57.8 114	101	91	. 1	127	139		114	174

a References:

- E. H. Amagat. 820 Values have been selected from his more extended Α table.
- E. H. Amagat. 320 As for A, but from another of his tables. P. W. Bridgman. Values computed from his specific volumes as given В in Table 95, Section III. T. W. Richards and W. N. Stull. 321
- RS
- RY
- W. Ramsay and S. Young.³²²
 L. B. Smith and F. G. Keyes.²⁹⁴ Values derived directly from their table SK of specific volumes.
- G. Tammann and W. Jellinghaus. Values computed from their specific volumes as given in Table 95, Section V; the unit of pressure is in this case 1 kg*/cm², but the difference between that and 1 atm is of little TI importance here.

Table 107.—Adiabatic Compressibility of Water

(For natural waters, see Table 110.)

The most accurate available values for the adiabatic compressibility, β_a , of water are probably (R), those derived by C. R. Randall ³¹⁷ from his

²⁰⁰ Amagat, E. H., Ann. de chim. et phys., (6) 29, 68-136, 504-574 (548, 549) (1893).

²²¹ Richards, T. W., and Stull, W. N., Carnegie Inst. of Washington, Publ. No. 7 (1903) → J. Am. Chem. Soc., 26, 399-412 (1904), and Z. physik. Chem., 49, 1-14 (1904).

⁸³² Ramsay, W., and Young, S., Phil. Trans. (A), 183, 107-130 (1892).

Table 107—(Continued)

measurements of the velocity of supersonic vibrations ($\beta_a = 1/V^2 \rho$, V =velocity, ρ = density). Earlier determinations (HL) by the same method were made by J. C. Hubbard and A. L. Loomis, 322a and similar determinations (P), based on the velocity of waves of audio-frequency, have been made by L. G. Pooler.³²³ D. Tyrer ³¹⁸ determined β_a directly (T) from the observed expansion that accompanied a sudden reduction of pressure from 2 atm to 1 atm. These 4 sets of data (R, HL, P, and T) are given below.

J. Claeys, J. Errera, and H. Sack 324 have suggested that the adiabatic compressibility of water may exhibit a type of hysteresis.

Unit of $\beta_a = 1$	cm² per megadyne = 1.01325	per atm.	Temp. $= t$ °C.
-----------------------	----------------------------	----------	-----------------

Method→	Veloci	ity	Dire	ect
Source ^a →	R	HL 1068	P	T
0 5	50.77	50.53 49.06		49.59
10 15	47.71	47.66 ^b 46.47		47.13°
20	45.54	45.48ª		45.00
25 30 35	44.05	44.69° 44.05 43.51	45.42 44.72	43.52
40 50	43.08 42.48	43.02	43.72 43.09	42.68 42.24
60 70 75	42.25 42.28		42.90 42.95 43.01	42.07 41.90
80	42.58		10.01	41.99
86 90 100	42.87			42.14 42.34

^a Source: See head-matter.

b. Colladon and C. Sturm ^{3.5} found $10^{\circ}\beta_{\alpha} = 49.5$ atm⁻¹ at 10° C, which must be reduced by 1.65, ^{3.50} giving $10^{\circ}\beta_{\alpha} = 47.8$ atm⁻¹ = 47.2 cm² per megadyne.

° R. W. Boyle, J. F. Lehmann, and S. C. Morgan ^{3.57} found from the velocity of supersonic waves that $10^{\circ}\beta_{\alpha} = 46.1$ cm²/megadyne at 12° C.

⁴ A. Pasuinskii ^{3.58} found $10^{\circ}\beta_{\alpha} = 45.5$ cm²/megadyne at 20° C.

[•] S. Parthasarathy 829 found $10^{\circ}\beta_a = 44.9 \text{ cm}^2/\text{megadyne}$ at 24 °C.

⁸²²a Hubbard, J. C., and Loomis, A. L., Phil. Mag. (7), 5, 1177-1190 (1928).

⁸²⁸ Pooler, L. G., Phys. Rev. (2), 35, 832-847 (1930).

⁸²⁴ Claeys, J., Errera, J., and Sack, H., Compt. rend., 202, 1493-1494 (1936).

⁸²⁵ Colladon, D., and Sturm, C., Mém. Sav. Etrang. Inst. Paris, 5, 267-347 (1838); Ann. de chim. et phys., 36, 113-159, 225-257 (1827); Ann. d. Physik (Pogg.), 12, 39-76, 161-197 (1828); "Mém. sur la compression des liquides et la vitesse de son dans l'eau," 1827; Ch. Schuchert, Genf,

⁸²⁰ Bungetzianu, D., Bull. Soc. Roumaine Sci. (Bucarest), 19, 1224-1246 (1229) (1910).

Boyle, R. W., Lehmann, J. F., and Morgan, S. C., Trans. Roy. Soc. Canada, III (3), 22, 371-378 (1928).

³²⁸ Pasuinskii, A., Acta Physiochim, URSS, 3, 779-782 (1935).

²⁰⁰ Parthasarathy, S., Proc. Indian Acad. Sci. (A), 2, 497-511 (1935).

⁸⁸⁰ Beattie, J. A., Int. Crit. Tables, 3, 100 (1928).

³³¹ Knudsen, M. H. C., "Hydrographische Tabellen," Copenhagen, 1901.

⁸⁸² Buchanan, J. Y., Proc. Roy. Soc. (London) (A), 23, 301-308 (1875).

⁸⁸⁸ Hill, E. G., Proc. Roy. Soc. Edinburgh, 27, 233-243 (1907).

Table 108.—Density of Sea-water: Pressure = 1 Atmosphere

(For density at temperature of maximum density, see Table 134.)

Sea-water contains about 35 g of salts per kg of sea-water. For the composition of the salts, and variations in the salinity and the temperature of sea-water, see Section 100.

The following data are derived from a table by J. A. Beattie,³³⁰ based primarily on the data given by M. H. C. Knudsen ³³¹ and J. Y. Buchanan,⁸³² but with a consideration of those of E. G. Hill,³³³ J. J. Manley,³³⁴ Dittman,³³⁵ F. L. Ekman,³³⁶ R. Lenz,³³⁷ C. O. Makaroff,³³⁸ C. von Neumann,³³⁹ and T. E. Thorpe and A. W. Rücker.³⁴⁰

Unit of $\rho = 1$ gram/ml; $(\rho = 1 + 10^{-5}\Delta)$; % = per cent by weight; salts = total salts. Temp. = l °C. Pressure = 1 atm.

				rressure =	1 atm.				
%C1	% salts	0	5	10	15	Δ	25	30	35
0.1	0.184	140	149	120	58	- 34	-151	-292	-455
0.2	0.364	287	293	261	197	+104	- 15	-158	-322
0.3	0.545	433	436	402	335	241	+120	-24	-189
0.4	0.725	579	579	542	474	377	256	+110	- 56
0.5	0.906	725	722	683	612	514	391	245	+ 77
0.6	1.086	871	865	823	751	651	526	379	210
0.7	1.267	1016	100 <i>7</i>	963	889	787	661	513	343
08	1.447	1162	1150	1103	1027	924	796	647	476
0.9	1.628	1307	1292	1243	1165	1060	931	780	608
1.0	1.808	1452	1434	1383	1303	1196	1066	914	741
1.1	1.989	1597	1577	1523	1441	1333	1201	1048	874
1.2	2.169	1742	1719	1663	1579	1469	1336	1182	1007
1.3	2.350	188 <i>7</i>	1861	1803	1717	1605	1472	1316	1140
1.4	2.530	2032	2003	1943	1855	1742	1607	1450	1274
1.5	2.711	2177	2146	2083	1993	1879	1742	1585	1407
1.6	2.891	2322	2288	2223	2131	2016	1878	1720	1541
1.7	3.072	2468	2431	2364	2270	2153	2014	1855	1675
1.8	3.252	2613	2574	2504	2408	2290	2150	1989	1809
1.9	3.433	2758	2716	2644	2547	2427	2286	2124	1944
2.0	3.613	2904	2859	2785	2686	2564	2422	2260	2079
2.1	3.794	3049	3002	2926	2825	2701	2558	2395	2214
2.2	3.974	3195	3145	3067	2964	2839	2695	2531	2349
2.3	4.155	3341	3289	3208	3104	2978	2831	2667	2484
							_504	_50,	0 .

Table 109.—Specific Volume of Sea-water: Pressure Greater than 1 Atm (For the specific volume at 1 atm, take the reciprocal of the density as given in Table 108.)

The data in Section A of this table, taken from the compilation of L. H. Adams,⁸⁴¹ are from V. Bjerknes and J. W. Sandström ⁸⁴² and are

⁸⁸⁴ Manley, J. J., Idem, 27, 210-232 (1907).

²⁸⁵ Dittman, "Rep. Sci. Results, Physics and Chemistry, Voyage H. M. S. Challenger," Vol. 1, 1889.

³³⁶ Ekman, F. L., Kongl. Svenska Vet-akad. Handl., 9, No. 4, 1870.

⁸⁸⁷ Lenz, R., Mém. de l'acad Sci. Russie (7), 29, No. 4, (1881) → Fortschr. d. Physik, 38₂, 661-662 (1888).

²⁸⁸ Makaroff, C. O., J. Russ. Phys. Chem. Soc. (Chem.), 23, II, 30-88 (1891).

⁸⁰⁰ von Neumann, C., Ann. d. Physik (Pogg.), 113, 382 (1861).

⁸⁴⁰ Thorpe, T. E., and Rücker, A. W., Phil. Trans. (A), 166, 405-420 (1876).

Table 109—(Continued)

based upon the observations of V. W. Ekman.³⁴³ Differences of successive values of Δ are printed in distinctive type; in the first two subsections they are between the values from which they have been derived, in the third subsection they are to the right of the greater Δ .

Those in Section B are from the detailed practical tables of N. H. Heck and J. H. Service ³⁴⁴ based upon the very extensive tables appended to the publication by Bjerknes and Sandström already mentioned. The values here tabulated were taken directly from the tables of Heck and Service; those they give for other salinities and temperatures may be reproduced, usually within 2 units in the 5th significant digit, by means of the formula

$$\Delta_{s,t,d} = \Delta_{35,0,d} - (6.48 + 0.00375d)t - 0.46t^2 + (\delta - 0.283t - 0.005t^2) \cdot (s - 35)$$

Within the limits they consider $(s = 31 \text{ to } 37, t = 0 ^{\circ}\text{C} \text{ to } t_m, t_m \text{ being here tabulated})$, δ varies with d as shown in the final portion of the table. The units are as indicated below.

Salinity = s = total salts per kg of sea-water; specific volume = v_e = $1 - 10^{-5}\Delta$; pressure = P; depth below surface = d = $(d_1 + d_2)$; temperature = t °C; $\Delta_{e,t,d}$ = value of Δ for salinity s, temperature t, and depth d; $\Delta_{35,0,d}$ = value of Δ for s = 35 g/kg, t = 0 °C, depth = d; P_d = pressure at depth d.

Unit of P=1 bar = 0.9869 atm; of s=1 g/kg; of $v_0=1$ ml/g; of d=1 fathom = 6 ft = 182.88 cm.

				Sect	ion A				
$t \rightarrow P$	0		4.97		9.97		14.96		19.96
(s=31.130)									
0	2440 888	35	2405 864	65	2340 <i>845</i>	89	2251 830	113	2138 <i>820</i>
200	3328 <i>832</i>	59	3269 812	84	3185 796	104	3081 783	123	2958 772
400	4160 784	79	4081 766	100	3981 751	117	3864 739	134	3730 731
600	4944	97	4847	115	4732	129	4603	142	4461
(s = 38.525)									
0	3004 <i>866</i>	45	2959 845	72	288 7 830	96	2791 813	117	2674 803
200	3870 <i>813</i>	66	3804 793	87	371 7 775	113	3604 767	127	3477 757
400	4683 766	86	4597 <i>74</i> 9	105	4492 736	121	4371 725	137	4234 717
600	5449	103	5346	118	5228	132	5096	· 145	4951

⁸⁴¹ Adams, L. H., Int. Crit. Tables, 3, 439-440 (1928).

⁸⁴² Bjerknes, V., and Sandström, J. W., Carnegie Inst. Washington, Publ. 88, Part I (1910).

²⁴² Ekman, V. W., Conseil. perm. int. l'explor, de la Mer, Publ. de Circon. No. 43, Copenhagen, 1908.

³⁴⁴ Heck, N. H., and Service, J. H., U. S. Coast and Geod. Survey, Spec. Publ. No. 108, (1924).

Table	109(Continued)	
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t=0 °C; s=3	35.00										
P		- Δ	_	P		_ A		P		_ A .	
0	2736			400	443			800	5940		361
100	3181	44.		500	484			900	6291		35 1
200	3612			600	520			1000	66 33		342
300	4030	41	8	700	557	79 370					
					Section	on B					
$d_1 \rightarrow$	100	300		500	700	900	100	300	500	700	900
d ₂				Δ.15,0,d					_ tm		
0	2736°	298		3147	3309	3469	22	22	22	20	12
1000	3627	378		3939	4093	4245	8	6	4	4	3
2000	4394	454		4692	4837	4983	3 2	3 2 2	3 2 2	4 3 2 2	3 2 2
3000	5126	527		5513	5553	5691	2	2	2	2	2
4000	5 828	596	5	6099	6234		2	2	2	2	
$d_1 \rightarrow d_2$	100	300	500 - δ -	700	900	100	300	500	7	00	900
0	76.3	75.3	75.0	74.3	73.8		55.4	92.3	129	3	166.5
1000	73.3	72.8	72.3	71.8	71.3	203.6	240.6	278.0	315		352.7
2000	70.8	70.3	69.8	69.5	69.0	390.2	427.5	465.2	502		540.4
3000	68.5	68.2	67.8	67.2	66.8		616.7	654.7	692		730.8
4000	66.2	66.0	65.8	65.2		768.8	807.0	845.1	883		

^a This value appears to be too great, the difference between the value for d=100 and d=300 being 249, while the other differences for 200 fathoms are about 160. In fact, the value 2736 is that for the surface (See P=0, end of Section A).

(Continued from p. 225.

M. Thiesen, K. Scheel, and II. Diesselhorst.²⁷⁴ Believing it better to keep the two sets distinct, I have not included that table in this compilation. The greatest difference between the two sets is 113 parts in 10⁷, and occurs at 41 °C.

Thiesen, Scheel, and Diesselhorst used the method of balanced columns of liquids, and their observations indicate that metal was dissolved from the tubes by the water. The amount so dissolved during the course of their second series of determinations was inferred, from the change in the electrical conductivity of the water, to have been such as to affect the density by 10 parts in a million. They endeavored to eliminate the effect of such solution by suitably combining related sets of data. Their observations were not very closely spaced with reference to the temperature. They combined their observations so as to obtain the density (ρ) at exactly 5° intervals from 0 to 40 °C. From these 9 values they determined the constants in the formula

$$1 - \rho = \frac{(t - 3.98)^2}{503570} \cdot \frac{t + 283}{t + 67.26}$$

and by means of that formula computed the values given in their table. Chappuis used the weight-thermometer method, using both glass and platinum-iridium bulbs; observations were made at many temperatures, closely spaced and well distributed. He represented them by a triad of

formulas in powers of the temperature (t °C), the constants being determined with high precision (see Table 99). His table was computed by means of these formulas. Quite recently it has been found 275 that there are systematic differences between the values in that table and the actual observations of Chappuis, and that the actual observations can be more closely represented by the single formula

$$1 - \rho = \frac{(t - 3.9863)^2}{508929.2} \cdot \frac{t + 288.9414}{t + 68.12963} \tag{1}$$

than by Chappuis's triad; and a table has been computed by means of that formula. That table is given in full in this compilation, together with the amounts by which each of the other two tables differ therefrom.

The data given by P. W. Bridgman in his compilation ²⁷⁶ are based upon those he had previously published, ²⁷⁷ and, with his recent paper, ²⁷⁸ are the source of most of the data here attributed to him.

An early study of the expansion of water in the range -13 to +100 °C was made by C. Despretz.²⁷⁹ but the compiler has not found those data; in the range -9 to +100 °C by the same investigator ²⁸⁰; and in the range -4 to -10 °R by Salm-Horstmar.²⁸¹

Tables and charts for compressed water, based, in the main, on data obtained at the National Bureau of Standards and at the Massachusetts Institute of Technology, have been published by J. H. Keenan.²⁸²

Dissolved Air: Effect on Density.

Dissolved air decreases the density of water. The frequently quoted differences reported by W. J. Marek 283 are the differences between the densities of water that has been freed from air by exhaustion just before measurement and those of water, at the same temperatures, that has merely been exposed to air for intervals of 1 to 3 days.²⁸⁴ They refer to the rather ill-defined conditions generally encountered in practice, rather than to the extreme conditions of complete air-freedom and air-saturation of Chappuis' work. Furthermore, when they are applied to those values for the density of air-saturated water which are published in the same paper, they lead to values of the density quite different from those generally accepted for airfree water (Table 93). The paper does not contain sufficient details to enable one to determine either the cause of the discrepancy or the accuracy of the differences, which are, in fact, about 43 per cent smaller than those published in his earlier (1884) paper. The several reported results are shown in Table 92, p. 198. P. Chappuis ²⁸⁴ and W. A. Adeney, A. G. G. Leonard, and A. Richardson 286 have studied the aeration of quiescent water. The former found that at 13.5 °C a layer 12 cm below the surface became half saturated in about a day, and 3/4 saturated in something over 4 days.

Recently, H. J. Emeléus et al.²⁸⁷ have reported that saturating water at 20 °C with air reduces its density by an amount equal to that caused by increasing the temperature of the water by 0.01 °C; that is, by 2 in 10⁶. That is much greater than the value given in Table 92. On the other hand, T. W. Richards and G. W. Harris²⁸¹ found by essentially the same method that the change in density under those conditions is less than 2 in 10⁷, which is about one-half the tabulated value.

Isothermal Compressibility of Natural Water.

Three sets of data are given in Table 110. The first two have ultimately been either taken or derived from the tables given by V. Bjerknes and J. W. Sandström³⁴² which in turn are based upon the observations of V. W. Ekman.³⁴³ The first was selected by L. H. Adams ³⁴¹ for his compilation and gives β for each of several pressures. The second has been adapted from the detailed practical tables compiled by N. H. Heck and J. H. Service ³⁴⁴ from the tables of Bjerknes and Sandström. It gives the values of the compression ($\beta' \equiv -(dv^*/dp)_t$, $v^* =$ specific volume) at 0 °C, salinity = 35 g/kg, for each of several depths below the surface. If we write $\beta'_{s,t,a}$ for the value of β' for sea-water of salinity s, temperature t, and depth d, then the values that Heck and Service give for the ranges s = 31 to 37, t = 0 to t_m may be reproduced very closely by means of the formula

$$10^{5}\beta'_{s,t,a} = 10^{5}\beta'_{35,0,a} - (0.0238 - 2(10^{-6})d)t + 0.000312t^{2} - (0.015 - 1.1(10^{-6})d) \cdot (s - 35)$$

the units being those named below. The pressures corresponding to the several values of d may be found in Table 109.

The third set covers earlier data frequently quoted but less reliable than those covered by the other two. They were obtained by P. G. Tait ³⁴⁵ and supersede those published in his earlier papers. ³⁴⁶ He summarizes these data in the following formulas:

Spring water:

$$10^{7}\beta_{m} = 520 - 17p + p^{2} - (355 + 5p)\frac{t}{100} + (3 + p)\frac{t^{2}}{100} \text{ per atm, and}$$

$$10^{5}\beta_{m} = \frac{186}{36 + p} \left(1 - \frac{3t}{400} + \frac{t^{2}}{10000}\right) \text{ per atm.}$$

Sea-water:

$$10^5 \beta_m = \frac{179}{38 + f} \left(1 - \frac{t}{150} + \frac{t^2}{10000} \right)$$
 per atm, and at 0 °C $10^5 \beta_m = 481 - 21.25 p + 2.25 p^2$ per atm.

For solutions of NaCl at 0 °C, he gives:

 $10^5\beta_m = 186/(36 + p + s')$ per atm, when the solution contains s'

grams of NaCl per 100 grams of water; s' was varied from 3.88 to 17.63. In neither pair are the two equations identical, but each was supposed to represent the data satisfactorily. In each of these equations, β_m is the mean compressibility between 1 atm and the pressure of p (long) tons* per sq. in.

The sea-water was not more particularly described; neither the composition nor the density is stated. Both it and the spring water were, presumably, nearly saturated with air, though nothing seems to have been said about this in the original articles. In the last section of the table certain values of β_m for pure, air-free water (from Table 105) are given together with the corresponding ones as derived from Tait's equations.

Symbols:
$$\beta = -\frac{1}{v_1} \left(\frac{\delta v}{\delta p} \right)_t$$
; $\beta_m = +\frac{1}{v_1} \left(\frac{v_1 - v}{p - A} \right)_t$; $\beta' = -\left(\frac{\delta v}{\delta p} \right)_t$; $v = -\left(\frac{\delta v}{\delta p} \right)_t$;

specific volume; p = pressure; A is value of p corresponding to 1 atm; v_1 is value of v at t °C and a pressure of 1 atm; the value of the pressure may be represented by P, P_b , or p, or indicated by b; b = depth below the surface of the ocean; s = salinity; $t_m = \text{highest temperature for which Heck and Service give data against which the formula given above for <math>\beta'_{s,t,d}$ may be checked.

Table 110.—Isothermal Compressibility of Natural Waters

(For source of data, explanation of symbols, etc. see text.)

Unit of $P_b = 1$ bar = 0.9869 atm; of P = 1 atm; of p = 1 (long) ton*/in²; of $\beta = 10^{-6}$ per bar; of $\beta_m = 10^{-6}$ per atm; of $\beta' = 1$ (cm³/g) per bar; of $d = (d_1 + d_2) = 1$ fathom = 6 ft = 182.88 cm; of s = 1 g/kg. Temp. = t °C.

A. Sea-water, 0 °C, 35 g salts per kg of sea-water.

B. Sca-water, 0 °C, 35 g salts per kg of sea-water.
$$\beta' \equiv \beta'_{35,0,d}$$
, $\beta' = (\delta v/\delta p)_t$
 $d_1 \rightarrow 0$ 300 500 700 900 100 300 500 700 900
0 4.50 4.44 4.39 4.33 4.28 22 22 22 20 12
1000 4.24 4.19 4.14 4.09 4.03 8 6 4 4 3
2000 4.00 3.94 3.91 3.87 3.83 3 3 3 3 2
3000 3.78 3.73 3.70 3.65 3.62 2 2 2 2 2 2 2
4000 3.58 3.55 3.52 3.47 2 2 2 2 2 2

C.
$$\beta_m \equiv (v_1 - v_p)/v_1(p-1)$$
; $10^6 \beta_m = a + bt + ct^2$.

Water→			- Spring -		Sea					
Þ	P	a	-b	c `	a	- b	c			
0	1	52.0	0.355	0.003	48.1	0.340	0.003			
1	152	50.4	0.360	0.004	46.2	0.320	0.004			
2	305	49.0	0.365	0.005	44.8	0.305	0.005			
3	457	47.8	0.370	0.006	43.8	0.295	0.005			

Rep. Sci. Results Voy. H. M. S. Challenger, Phys. and Chem.," Vol. 2, Part 4, London, Edinburgh, and Dublin, 1888.
 Tait, P. G., Proc. Roy. Soc. Edinburgh, 12, 223-224, 757-758 (1884); 15, 84 (1887).

$t\rightarrow$		0			10			20	
Watera→	Spr.	Pure	Sea	Spr.	Pure	Sea	Spr.	Pure	Sea
1	52.0	51.5	48.1	48.7	48.4	45.0	46.1	46.4	42.5
150 300	50.4 49.0	50.5	46.2	47.2	47.6	43.4	44.8 43.7	46.0 44.6	41.4 40.7
450	49.0 47.8	49.2 48.0	44.8 43.8	45.8 44.7	46.4 45.3	42.3 41.4	42.8	43.6	39.9

Table 110—(Continued)

The 3 samples of water are spring (= Spr.), pure air-free (= Pure), and seawater (= Sea). The data for the first and third have been computed by means of Tait's equations; those for the pure water have been taken from Table 105.

Adiabatic Compressibility (β_a) of Natural Waters.

From the observed velocity of sound generated by explosions in the sea, A. B. Wood, H. E. Browne, and C. Cochrane ³⁴⁷ have concluded that, for sea-water at 16.95 °C, under a mean pressure of 2 bars, and containing 35 g of salts per kg of sea-water, $10^6\beta_a = 42.7$ per bar = 43.3 per atm. As usual, $\beta_a \equiv -\frac{1}{v} \left(\frac{\delta v}{\delta P} \right)_c$.

33. MECHANICAL EQUIVALENT OF THE CALORIE

By the mechanical equivalent of the calorie is meant the work required to produce the amount of heat designated as one calorie.

Several different calories have been used and must be distinguished if a higher accuracy than 1 or 2 in 1000 is desired. For this reason, among others, it is desirable to express quantities of heat in terms of a less ambiguous unit, such as the joule. For uncertainties in the value of the international joule, see Section 2.

Of the various calories that have been used, four are of particular importance, having received widespread recognition. They are designated and defined thus: $1 \text{ cal}_{15} = \text{amount of heat required to raise 1 gm of water from 14.5 to 15.5 °C; 1 cal₂₀ = amount of heat required to raise 1 gram of water from 19.5 to 20.5 °C; 1 cal_m = 1 mean calorie = 1/100 of the amount of heat required to raise 1 gram of water from 0 to 100 °C, and 1 cal (ST) = 1 cal (steam) = I Int. cal. = 1 steam-table calorie = 1/1000 of the heat that is equivalent to 1/860 international kilowatt-hour = 4.18605 Int. joules. For the first three, the water is to be under an air pressure of 1 atmosphere. The fourth, independent of the properties of any particular substance, was defined by the International Steam-Table Conference, meeting in London in 1929.³⁴⁸$

A fifth calorie (cal_{ms}), a mean calorie based upon air-free saturated water, has been proposed by N. S. Osborne, H. F. Stimson, and E. F. Fiock ³⁴⁹ and defined as 1/100 of the change in the enthalpy ("heat con-

⁸⁴⁷ Wood, A. B., Browne, H. E., and Cochrane, C., Proc. Roy. Soc. (London) (A), 103, 284-303 (1923).

Engineering (London), 128, 751-752 (1929); Z. Ver. deuts. Ing., 73, 1856-1858 (1929); Mech. Eng., 52, 120-122 (1930).
 Osborne, N. S., Stimson, H. F., and Fiock, E. F., Mech. Eng., 50, 152-153 (1928).

tent") of 1 gram of saturated water on passing from 0 to 100 °C. exceeds 1 cal_m by only about 0.001 joule.

R. Jessel 850 has held that the heat capacity of water is significantly affected by the presence of dissolved air, and that air-free water must be used if highly reproducible results are to be obtained. With that view T. H. Laby and E. O. Hercus 351 disagree. See also Table 112.

The various results obtained for the mechanical equivalent of the calorie have been reviewed and discussed by J. S. Ames, \$52 E. H. Griffiths. \$53

Table 111.—Mechanical Equivalent of the Calorie

The first value for cal₂₀ and for cal_m has been taken directly from the compilation by T. H. Laby and E. O. Hercus. 359

Unit of work = 1 joule = 107 ergs, unless value is followed by I (= Int. joule) calms. Ref.a calıs cal₂₀ ICT accepted ICT mean LH (1927) OSF (1928) OSG (1937) RTB (1929) 4.185b 4.186 4.181 4.1853 4.1818 4.1809 4.1876(I) 4.188 4.1876(I) 4.1852 HJ (1926) 4.1863 4.182 L (1933) 4.186

" References:

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erences:

"ICT accepted" are values accepted by the International Critical Tables [1, 18 (1926)].

"ICT mean" are the means given by Lahy and Hercus, and are based upon the work of: Barnes, H. T., Proc. Rov. Soc. (London) (A), 82, 390-395 (1909).

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Callendar, H. L., Idem, 212, 1-32 (1912).

Day, W. S., Phil. Mag. (5), 46, 1-29 (1898).

Griffiths, E. H., Proc. Roy. Soc. (London), 55, 23-26 (1894); Phil. Trans. (A), 184, 361-504 (1894); Idem, 186, 261-341 (1895).

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Rispail, L., Ann. de chim. et phys. (8), 20, 417-432 (1910).

Rowland, H. A., Proc. Amer. Acad. Arts Sci., 15, 75-200 (1880).

Schuster, A., and Gannon, W., Phil. Trans. (A), 186, 415-467 (1895).

Sutton, T. C., Phil. Mag. (6), 35, 27-29 (1918).

L.H. Laby, T. H., and Hercus, E. O., Phil. Trans. (A), 227, 63-92 (1927).
                                                                 Laby, T. H., and Hercus, E. O., Phil. Trans. (A), 227, 63-92 (1927).
Oshorne, N. S., Stimson, H. F., and Fiock, E. F. 340
Oshorne, N. S., Stimson, H. F., and Ginnings, D. C., J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937).
Value derived by R. T. Birge 350 from the observations of others.
The Reichsanstalt value as given by F. Henning and W. Jaeger, "Handb. d. Physik" (Scheel), Vol. 2, pp. 487-518 (497), 1926.
Conclusion of V. S. Lipine. 355
                      LH
                      OSG
                      RTB
                      НJ
<sup>b</sup> The value accepted for this compilation: 1 cal<sub>15</sub> = 4.185 joules.
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⁸⁵⁰ Jessel, R., Proc. Phys. Soc. (London), 46, 747-763 (1934).

AB1 Laby, T. H., and Hercus, E. O., Idem, 47, 1003-1008-1011 (1935). See also Hercus, E. O., Idem, 48, 282-284 (1936).
 Ames, J. S., Rapports Cong. Int. Phys. (Paris), 1, 178-213 (1900).

⁸⁵⁸ Griffiths, E. H., "Dictionary of Applied Physics" (Glazebrook), Vol. 1, pp. 477-494 (1922).

⁸⁵⁴ Laby, T. H., Proc. Phys. Soc. (London), 38, 169-172-175 (1926).

²⁸⁵ Lipine, V. S., Mém. prés à la VIII Conf. Gén. des Poids et Mes., 1933.

⁸⁵⁶ Birge, R. T., Rev. Mod. Phys., 1, 1-73 (1929).

ast Henning, F., and Jaeger, W., "Handb. d. Physik" (Scheel), Vol. 2, pp. 487-518, 1926). 858 Fiock, E. F., Bur. Stand. J. Res., 5, 481-505 (RP210) (1930) = Mech. Eng., 52, 231-242 (FSP-52-30) (1930).

859 Laby, T. H., and Hercus, E. O., Int. Crit. Tables, 5, 78 (1929).

T. H. Laby,³⁵⁴ and V. S. Lipine.³⁵⁵ R. T. Birge ⁸⁵⁶ has discussed them with special reference to the actual values of the standards used; and F. Henning and W. Jaeger,³⁵⁷ concluding that all determinations, except those made at the Physikalisch-Technischen Reichsanstalt, are vitiated by uncertainties regarding the actual values of the standards employed, rejected all except the Reichsanstalt's.

The several determinations of various thermal properties of saturated water have been reviewed by E. F. Fiock.³⁵⁸

34. THERMAL ENERGY OF WATER

In this section are considered the specific heat (c and C), the enthalpy or heat content $(H, H = E + pv, (\delta H/\delta t)_p = c_p)$, the "free energy at constant pressure" (the Gibbs function, $G, G = H - ST, G_T - G_{T0} = C_T$)

$$-T\int_{T_0}^T (H/T^2)dT$$
), the entropy (S), the heat of isothermal compres-

sion (Q), the decrease in the internal energy on isothermal compression (D), the increase in temperature on adiabatic compression (Joule-Thomson effect), and certain related quantities, all intimately related to the thermal energy of water.

Data referring to the thermal energy of water have been reviewed and discussed by E. F. Fiock ³⁵⁸ and by M. Jakob. ³⁶⁰

There are no direct determinations of the values of the specific heat of water at constant volume, of the ratios of the specific heats, or of their differences, but all of these can be computed from the observed compressibility, thermal expansion, and specific heat at constant pressure. Values so determined may be called *static* values. They can also be determined from the velocity of sound, in which case they may be described as *dynamic*. Likewise, the increase in temperature on adiabatic compression may be determined either statically, from the thermal expansion, or dynamically, from the observed drop in temperature that accompanies a sudden release of pressure.

If water consisted of a single species of molecule and if the internal state of a molecule were unaffected by gross dynamic changes in the substance, then no difference between the static and the dynamic values of those various thermal quantities would be expected. But there are reasons for believing that water may contain associated molecules of more than one type, and there is evidence indicating that the internal state of a molecule may be affected by gross dynamic changes in the substance. In which cases the static and the dynamic values of those thermal properties would be expected to differ, unless the times required to reëstablish equilibrium between the several types of molecules and between each type of molecule and the gross dynamic state of the substance are each negligibly short as

³⁶⁰ Jakob, M., Engineering (London), 132, 518-521, 550-551 (1931).

Table 112.—Effect of Dissolved Air on the Specific Heat of Water

R. Jessel 850 has stated that the presence of dissolved air increases the specific heat of water and lowers the temperature at which the minimum occurs; he presents the two sets of values of c here tabulated. He suggests that the calorie should be defined in terms of air-free water.

Laby and Hercus do not accept Jessel's conclusions; they conclude from thermodynamic considerations that the presence of dissolved air produces a negligibly small effect, and suggest that Jessel's observations may be explained by irregularities caused by escaping air-bubbles, as remarked by them in 1927.³⁶⁹ In the discussion following the Laby-Hercus paper, Jessel maintains his position, and develops his view of the subject.

The compiler has determined, and tabulated below, the excess of each of Jessel's values of c above that of the corresponding number (n) defined

by the formula $n = 4.185 \left[1 - 0.233 \left(\frac{t - 15}{1000} \right) + 6.32 \left(\frac{t - 15}{1000} \right)^2 \right]$; if $n \equiv 20 \, ^{\circ}$ C, n lies between the two sets of c.

		of $c = 1$ joule/g-°C	= 10 ⁷ ergs/g⋅°C.		
t	Ordinary	c - n	1	De-aerated -	c-n
12.3	4.1944	0.0066	16.8	4.1858	+0.0025
20.1	4.1857	0.0050	22.1	4.1780	-0.0014
25.2	4.1861	0.0083	32.9	4.1740	-0.0020
32.3	4.1890	0.0126	38.6	4.1732	-0.0035
37.1	4.1893	0.0129	43.7	4.1724	-0.0064
37.7	4.1892	0.0127	43.9	4.1736	-0.0053
42.5	4.1924	0.0142	50.2	4.1776	-0.0058
49.0	4.1916	0.0092	52.5	4.1773	-0.0083
			59.5	4.1772	-0.0168
			69.8	4.1845	-0.0265

Table 113.—Specific Heat of Compressed Water at 1 Atm or at Constant Volume

(For more highly compressed water see Table 115; for the limiting value as saturation is approached see Table 116; for sea-water see Table 129.)

The O values are the most accurate at present available. They are given directly; the others, by the amount (Δ) that each exceeds the corresponding O one.

Example: At 5 °C the A value is 4.20137 + 0.004 = 4.205, the JS value is 4.20137 - 0.004 = 4.197.

ant Tammann, G., and Ellmächter, A., Z. anorg. allgem. Chem., 200, 153-167 (1931).

⁸⁰² Awhery, J. H., Int Crit Tables, 5, 113 (1929).

⁸⁶⁸ Randall, M., Idem, 7, 232 (1930).

³⁶⁴ Jacker, W., and v. Steinwehr, H., Ann. d. Physik (4), 64, 305-366 (1921) → Sitz. d. K. preuss. Akad. Wiss. (Berlin), 1915, 424-432 (1915).

³⁰⁵ Osborne, N. S., Stimson, H. F., and Fiock, E. F., Mech. Eng., 51, 125-127 (1929).

⁸⁰⁰ Hofbauer, P. H., Atti pont. acc. sci. nouvi Lincei, 84, 353-363, 581-586 (1931).

²⁰⁷⁷ Koch, W., Forsch. Gebietc, Ingenieurw., 5, 138-145 (1934) → Z. Ver. deut. Ing., 78, 1110 (1934)

<sup>(1934).

***</sup>Bavliček, J., and Miškovský, L., Helv. Phys. Acta, 9, 161-207 (1936).

***Laby, T. H., and Hercus, E. O., Proc. Phys. Soc. (London), 47, 1003-1008-1011 (1935);

**Hercus, E. O., Idem, 48, 282-284 (1936).

Table 113—(Continued)

The data in Awbery's compilation are based upon $c_n = 4.190$ joules/ g.°C at 15 °C; those in Randall's on 4.182; those of Jaeger and v. Steinwehr on 4.1842. Here they have all been reduced to the same basis by multiplying the respective values of the ratio $(c/c_{15})_p$ by 4.185, the value, in absolute joules, accepted by the International Critical Tables. As 4.185 Int. joules is essentially the value at 15 °C found by O, all the values in the table may be regarded as expressed in Int. joules.

The values given for c_n have been derived by the compiler from the O values of c_p by the R and HL values given in Table 119.

Unit of cn. cr.	and $\Lambda = 1$ Int	inule/g.°C	Temp. = t °C	(Int. scale)

Ref. →	$c_p = 1$	A	R 1000 Δ. ——	JS	Co
	4.21753	-3	+5		4.2151
5	4.20137	+4	+5	-4	4.2012
0 5 10 15	4.19107	+4 +3	+2	-1	4.1865
15	4.18463	0	0	0	4.1706
20	4.18073	-2	-1	0	4.1535
25	4.17856		-2	0	4.1348
30	4.17751		$-2 \\ -4$	-2	4.1147
35	4.17734		-4	-1	4.0939
40	4.17772	-3	-4 -5	-1	4.0729
45	4.17860			+ 1	
50	4.17990		-5	+ 3	4.015
55	4.18153				
60	4.18354	-3	-4		3.976
65	4.18592				
70°	4.18873		-4		3.923
75	4.19191				
80	4.19551	-9	-3		3.852
85	4.19957				
90	4.20418		-2		3.790
95	4.20932				
100	4.21510	-22	-5		3.757

* References:

JS

^b A. Romberg ^{300a} has reported $c_{73} = (1.0040 \pm 0.0005) c_{20}$, whence $c_{73} = 4.197$, a very high value.

Awbery, J. H., 202 based upon the consideration of the work of Barnes, H. T., Proc. Roy. Soc. (London) (4), 67, 238-244 (1900); Phil. Trans. (4), 199, 149-263 (1902); Trans. Roy. Soc. Canada III (3), 3.27 (1909); Reports B. A. A. S., 1909, 403-404 (1909); Barnes, H. T., and Cooke, H. L., Phys. Rev., 15, 65-72 (1902); Bousfield, W. R., Proc. Roy. Soc. (London) (A), 93, 587-591 (1917); Bousfield, W. R. and W. E., Phil. Trans. (4), 211, 199-251 (1911); Callendar, H. L., Idem, 212, 1-32 (1912); Proc. Roy. Soc. (London) (A), 86, 254-257 (1912); Cotty, M. A., Ann. de chim. et phys. (8), 24, 282-288 (1911); Dieterici, C., Ann. d. Physik (4), 16, 593-620 (1905); Griffiths, E. H., Phil. Trans. (4), 184, 361-504 (1894); "Thermal Measurement of Energy," Cambridge Univ. Press, 1901; "Dictionary of Applied Physics" (Glazchrook's), vol. 1, pp. 477-494, London, Macmillan, 1922; Guillaume, C. E., Compt. rend., 154, 1483-1488 (1912); Jaeger, W., and v. Steinwehr, H., Ann. d. Physik (4), 64, 305-366 (1921); Ianke, Diss. Rostock (1910); Lüdin, E., Mitt. Naturw. Ges. Winterthur, Heft 2 (1900); Pagliani, S., Nuovo Cim. (6), 8, 157-188 (1914); Regnault, Mém. Acad. Roy. Sci. Inst. France, 21, 729-748 (1847); Rowland, H. A., Proc. Amer. Acad. Arts Sci., 15, 75-200 (1880).
Jaeger, W., and v. Steinwehr, H., 204 included in Würmetabellen (1919) compiled by L. Holborn, K. Scheel, and F. Henning. Osborne, N. S., private communication, 1938.
Randall, M., Int. Crit. Tables, 7, 232 (1930), based on work of Barnes, H. T., Phil. Trans. (A), 199, 159-263 (1902).

³⁰⁰a Romberg, A., Proc. Am. Acad. Arts Sci., 57, 375-387 (1922).

Table 114.—Mean Specific Heat of Water at 1 Atm 370

 $(\hat{c}_p = \text{mean specific heat between } t_1 \text{ and } t_2)$ Unit of $c_n = 1$ Int. joule/g.°C. Temp. t_1 and t_2 on Int. Centigrade scale. $t_1 \rightarrow t_2$ 25 50 55 60 4.17889 4.17826 4.18174 4.17998 4.17801 4.18006 910 860 847 161 170 948 910 906 027 65 70 977 195 4.18002 973 069 4.18061 237 124 4.18048 065 75 291 195 149 136 157

Table 115.—Specific Heat of Compressed Water at Constant Pressure (For values at a pressure of 1 atm see Table 113.)

If c_p and c_{p_1} are the specific heats at the constant pressures p and p_1 , respectively, and for the same temperature, then Δ_{p_1} is defined by the relation $c_p = c_{p_1} (1 + \Delta_{p_1})$. Values of c_{p_1} for each temperature and of $1000\Delta_{p_1}$ for each temperature and pressure are tabulated. Example: From Section I (Koch) we find for 260 °C and p = 300, $c_p = 4.944 (1 - 0.057) = 4.944 - 0.282 = 4.662$; likewise for 260 °C and p = 50, $c_p = 4.663 (1 + 0.060) = 4.663 + 0.280 = 4.943$.

Unit of $p = 1 \text{ kg*/cm}^2 = 0.9678 \text{ atm}$; of $c_p = 1 \text{ joule/g °C}$. Temp. = t °C

I. W. Koch.³⁶⁷ Smoothed values based on thermal determinations; precision does not exceed 0.004 joules/g·°C; conversion from Int. steam cal. to joules by the compiler; 1 cal = 4.186 joules.

$t \xrightarrow{t}$	50	100	150	200	250	300
	C60			$-1000\Delta_{50}$		4.0
0	4.203	2	4	6	8	10
20	4.169	2	4	7	9	12
40	4.161	2	5	8	10	13
60	4.165	2 2 2 3	6	9	12	13 15
80	4.182	4	4 5 6 7	10	14	17
100	4.203	4	7	11	15	18
120	4.232	4	8	12	16	20
140	4.266		10	13	17	22
160	4.324	4 5	10	15	19	24
180	4.395	6	11	17	22	28
200	4.483	7	13	20	26	32
220	4.592	8	16	23	30	37
240	4.738	10	19	28	37	45
260	4.944	13	25	36	46	57
$p \rightarrow t$	50	100	150 +1000Δ ₃₀₀ -	200	250	300
260	60	47	34	22	12	4.663
280	00	60	44	37 ,	14	4.860
300		106	63	35	16	5.119
310		100	91	48	20	5.262
			140	72	44	5.433
320						
330			223	110	44	5.659
340			361	175	73	5.965
350				278	102	6.430

⁸⁷⁰ Osborne, N. S., Private communication, 1938.

Table 115—(Continued)

II. M. Trautz and II. Steyer.³⁷¹ Computed from volumetric data and presented by small graphs; values read from graphs with a precision not exceeding 1 or 2 parts per 1000. Conversion to joules by the compiler.

$t \xrightarrow{t}$	50 C50	100	150	200 1000Δ ₅₀	250	300
0	4.169	1	5	6	9	10
5 0	4.169	4	6	8	11	14
100	4.203	4	8	10	15	17
150	4.282	6	13	18	22	28
200	4.479	10	19	24	34	42
250	4.998	20	31	45	59	74
t			+1000\Delta_200 -			C300
250	80	58	41	31	16	4.630
300			74	43	20	5.295
350				310	162	8.79

III. P. W. Bridgman.³⁷² Computed from volumetric data. Compiler scaled his small graph and converted the values from calories to joules. Precision not greater than 1 or 2 parts in 1000.

$c_1 \rightarrow$	0 4.21 ₈	20 4.18 ₅	40 4 18-	60 4.18 ₅	80 4.20 ₆
p			1000Δ ₁		
1000	+40	4 52	+55	+46	+30
2000	62	74	84	<i>7</i> 5	+18
3000	76	84	99	94	+2
4000	87	88	113	102	-12
5000	98	95	124	107	-25
5500	109	100	128	108	-31
6000	135	106	130	109	-38
7000		108	128	106	 53
8000		94	122	102	-70
9000			118	97	88
10000			114	90	-109
11000			108	83	-136
12000			100	75	-169

Table 116.—Specific Heat of Compressed Water: Limit as Temperature Approaches that of Saturation

The subscript " $t \to \text{sat}$ " is used to denote the limiting value approached as t approaches the temperature corresponding to saturation under the specified conditions. The value of $(c_v)_{t\to \text{sat}}$ may be obtained from that of $(c_p)_{t\to \text{sat}}$ by subtracting the corresponding value of (c_p-c_v) , given in Table 119.

Unit of $c_p = 1$ Int. joule/g. °C. Temp. = t °C (Int. scale)

- I. Preferred value at 100 °C is $(c_p)_{t\to sat} = 4.2151$ (see Table 113).
- II. SK^a Values computed by Smith and Keyes by means of an equation set up by them to represent several sets of data, including their own on the specific volume.

⁸⁷¹ Trautz, M., and Steyer, H., Forsch. Gebiete Ingenieurw., 2, 45-52 (1931).
872 Bridgman, P. W., Proc. Amer. Acad. Arts Sci., 48, 307-362 (Fig. 11) (1912).

Table 116—(Continued)						
ŧ	$(c_p)_{t \to sat}$	*	$(c_p)_{t \rightarrow uat}$	*	$(c_p)_{f \to sat}$	
0	(4.2208)	100	4.2127	190	4.4514	
1Ŏ	(4.1877)	100	4.2127	200	4.4958	
20	4.1772	110	4.2267	210	4.5465	
30	4.1747	120	4.2435	220	4.6066	
40	4.1763	130	4.2611	230	4.6755	
50	4.1765	140	4.2839	240	4.7560	
60	4.1808	150	4.3099	250	4.8423	
70	4.1869	160	4.3340	260	4.9651	
80	4.1930	170	4.3719			
90	4.2022	180	4.4137			

III. Miscellaneous values.

Ref.a→	0	SK	A	R
100	4.2151	4.2127	4.193	4.210
125 150		4.3099	4.218	4.235 4.265
200		4.4958	4.250	4.294
250		4.8423	4.29	
300			4.34	

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- Awhery, J. H. 362 For work considered by him, see Table 113, reference note and
- Awnery, J. R.—

 Osborne, N. S. 870

 Randall, M., Int. Crit. Tables, 7, 232 (1930), based on work of Barnes, H. T., Phil.

 Trans. (A). 199, 148-263 (1902). See head of Table 113.

 Smith, L. B., and Keyes, F. G., Proc. Amer. Acad. Arts Sci., 69, 285-314 (1934) →

 Mech. Eng., 56, 92 94 (1934). R

Table 117.—Specific Heat of Compressed Water at Constant Volume (Isopiestics)

Derived from a graph published by P. W. Bridgman 372, Fig. 12 and based upon his measurements of the compressibility. The values of c_v cannot be read more accurately than 1 or 2 parts in 1000.

In each case the volume is that corresponding to the indicated temperature and pressure. If c_1 and c_v = specific heat at constant volume for a pressure of 1 atm and of $p \text{ kg*/cm}^2$, respectively, the temperature being the same in each case, then $\Delta \equiv (c_1 - c_v)/c_1$ or $c_v = c_1(1 - \Delta)$. The values tabulated for c_1 correspond to those read from the graph. Example: At 40 °C and 3000 kg*/cm², $c_v = 4.07_2(1 - 0.124) = 3.56_7$ j/g·°C.

Unit of p=1 kg*/cm² = 0.9678 atm; of $c_1=1$ joule/g.°C. Temp. = t°C

			,	, 6	
$\begin{matrix} t \rightarrow \\ c_1 \rightarrow \end{matrix}$	0 4.22 ₃	20 4.14 ₇	40	60 3.96 ₇	80 3.85 ₀
P			1000A		
1000	49	62	63	56	16
2000	83	98	102	91	0
3000	108	119	124	113	-13
4000	128	131	147	129	-23
5000	143	139	167	140	-33
6000	162	162	183	148	-43
7000		195	193	154	-54
8000		211	200	159	-65
9000			205	161	-80
10000			209	163	-100

Table 117(Continued)
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$t \rightarrow c_1 \rightarrow c_2 \rightarrow c_1 \rightarrow c_2 \rightarrow $	0 4.22 ₃	20 4.14 ₇	40 4.07 ₂ 1000Δ	60 3.96 ₇	80 3.85 ₀
11000			209	161	-126
12000			205	155	-165

Table 118.—Specific Heat of Compressed Water at Constant Volume (Isometrics)

Derived from a graph published by P. W. Bridgman $^{372, \text{ Fig. } 13}$ and based upon his measurements of the compressibility. The values of c_v cannot be read more accurately than 1 or 2 parts in 1000.

If $c_{1,0} = c_v$ for $v^* = 1$ and t = 0 °C, then $\Delta \equiv (c_{1,0} - c_v)/c_{1,0}$; or, the specific heat at constant volume (c_v) for the volume and temperature indicated is $c_v = c_{1,0}(1 - \Delta)$; $v^* =$ specific volume. Bridgman gives $c_{1,0} = 1.000$ cal/g.°C = 4.185 joules/g.°C.

	Unit	of $v^* = 1 \text{ cm}^3/$	g. Temp. = to	C	
$t \rightarrow v^*$	0	20	40 1000Δ	60	80
1.025					84
1.000	0	12	43	80	97
0.975	22	50	83	112	91
0.950	47	85	116	140	75
0.925	76		140	160	62
0.900	105	133	166	176	50
0 875	126	145	193	190	39
0.850	150	188	216	202	17
0 825		219	230	206	-24
0 800			228		

compared with that for the change in pressure involved in the dynamic method.

In the case of the decrease of temperature on adiabatic expansion, G. Tammann and A. Elbrächter ³⁶¹ have sought evidence for such a difference. They have found differences, but have been unable to explain them in terms of the expected type of changes in the association (see Table 126).

Table 119.—Ratio and Difference of the Principal Specific Heats of Water: 1 Atm or Saturation

(For values at higher pressures, see Table 120; for sea-water, see note b.)

If c_p and c_v = specific heat of water at constant pressure and constant volume, respectively, $\gamma = c_p/c_v$; if $\alpha = (1/v) (dv/dt)_p$, v = volume of any fixed mass of water, T = absolute temperature, and V = velocity of sound, then $\gamma - 1 = T\alpha^2V^2/c_p$; $c_p - c_v = c_p(\gamma - 1)/\gamma$. Bridgman's values (B) for $(c_p - c_v)$ have been derived from his graphs, ³⁷², Figs. 11, 12 which are based upon his measurements of the compressibility.

The values at saturation are those derived from the limiting values

Table 119—(Continued)

approached by c_v and c_v as t approaches the temperature at which water is saturated at the coexisting values of p and v, respectively.

Unit of $(c_p - c_p) = 1$ Int. joule/g.°C = 0.23895 cal₁₅/g.°C. Temp. = t°C

I. Pressure = 1 atm = 1.03323 kg*/cm^2 .

Basis ^a →	R	S	HL	R	HL	В
ŧ	10	$4(\gamma-1)$	$\equiv \delta_{\gamma}$	10 ³ ($(c_p - c_v) \equiv \delta$	
0	5.81	5	5.85	2.45	2.46	
5			0.34		0.14	
0 5 10	10.84	20	10.86	4.53	4.54	
15			33.68		14.00	
20	65.5	60	65.6	27.20	27.23	30
25			105.8		43.8	
30	152.7	$14_{\rm o}$	152.5	62.8	62.7	
35			203.7		83.4	
40	257.5	27.	257.5	104.9	104.8	110
50	385	38		155		
60	524	62 ₀		208		220
70	676	- 80 ₀		266		•
80	840	1000		324		360
86	942			361		•
90	110°	110_{o}		414°		
100	123°	114_{o}		458°		

II. Values at saturation. Derived from the values of c_p and c_v as computed by Smith and Keyes (see Table 116). $\delta_{\gamma} \equiv 10^4 (\gamma - 1)$, $\delta_c \equiv 10^3 (c_p - c_v).$

ŧ	δ_{γ}	δ_c	t	δ_{γ}	δ_c	ŧ	δ_{γ}	δ_o
0	1.5	0.6	90	999	381.4	180	2903	993.9
10	13.4	5.6	100	1184	446.1	190	3268	1095.3
20	63.3	26.3	110	1381	512.7	200	3540	1175.5
30	143.5	59.1	120	1586	580.8	210	3831	1259.7
40	246.7	100.5	130	1808	652.1	220	4130	1346.2
50	369	148.5	140	2023	721.3	230	4440	1437.4
60	506	201.5	150	2257	793.9	240	4750	1531.9
70	658	258.7	160	2499	866.1	250	5120	1640.5
80	822	318.8	170	2743	941.7	260	5460	1752.4

- ^a Except as otherwise noted, the bases on which these values rest are:
 - C. R. Randall's determination of V^{sta} ; c_p from Table 113, Column A (or R if no value in A); $(1/v) (dv/dt)_p$ from equations of P. Chappuis, 0° to 40°, or of Thiesen 40° to 86° (Table 99); computation by the compiler. J. C. Hubbard and A. L. Loomis 374 from their own determinations of the velocity of sound; c_p from Table 113 (Column JS), Chappuis' equations. P. W. Bridgman, from his determinations of the compressibility (cf. Tables 115 and 117) R

 - В Tables 115 and 117).
 - F. A. Schulze. 875 Computed by him from isothermal compressibility S and c_p ; the same values are given in each paper.

For sea-water at 16.95 °C and pressure = 2 atm, salinity = 35 g/kg, $10^4(\gamma - 1)$ = 94 = 5, from velocity of sound and isothermal compressibility.**

Computed by D. Tryer **re from the isothermal compressibility and c_p.

⁸⁷⁸ Randall, C. R., Bur. Stand. J. Res., 8, 79-99 (RP402) (1932).

⁸⁷⁴ Hubbard, J. C., and Loomis, A. L., Phil. Mag. (7), 5, 1177-1190 (1928).

²⁷⁵ Schulze, F. A., Z. physik. Chem., 88, 490-505 (1914); Physik. Z., 26, 153-155 (1925).

⁸⁷⁰ Tyrer, D., J. Chem. Soc. (London), 103, 1675-1688 (1913); Z. physik. Chem., 87, 169-181 (1914).

Specific Heat of Water.

In the International Critical Tables, two sets of values for the specific heat of water at constant pressure (c_p) are given: those compiled by J. H. Awbery 362 and those by M. Randall. 363 The first are considered the more accurate. Neither agrees with the set published by W. Jaeger and H. v. Steinwehr 364 and included in the "Wärmetabellen" (Vieweg, Braunschweig, 1919) compiled by L. Holborn, K. Scheel, and F. Henning (see Table 113). A graphical comparison of the more important published values has been given by N. S. Osborne, H. F. Stimson, and E. F. Fiock. Among the various interpolation equations that have been proposed may be mentioned those by P. H. Hofbauer, 366 L. B. Smith and F. G. Keyes, 204 W. Koch, 367 and J. Havliček and L. Miškovský. 368 None of the earlier ones is satisfactory if more than moderate accuracy is desired.

Table 120.—Ratio and Difference of the Principal Specific Heats of Water under High Pressure

Derived from graphs constructed by P. W. Bridgman ³⁷², Figs. 11, 12 * from his measurements of the compressibility. The specific heats cannot be read from the graphs to a higher accuracy than 1 or 2 parts in 1000. The values of γ have been computed from those of $(c_p - c_v)$ as determined from the graphs.

<i>t</i> →	0	20	40	60	80	0	20	40	60	80
Þ		1	$000(\gamma -$	1) ——			10	0(cp - 1	cv) ——	_
0	0	7	27	55	93	0	3	11	22	36
1000	7	21	37	67	77	3	8	14	25	29
2000	23	37	46	72	73	9	14	17	26	28
3000	34	49	56	77	77	13	18	20	27	30
4000	46	61	69	87	81	17	22	24	30	32
5000	50	62	83	97	83	18	22	28	33	33
6000	31	78	93	104	85	11	27	31	35	34
7000		117	113	113	91		39	37	38	37
8000		159	127	126	98		52	41	42	40
9000			139	139	99			45	45	41
10000			152	148	102			49	49	43
11000			158	153	102			51	51	44
12000			163	155	96			53	52	43

Unit of $(c_p - c_v) = 1$ joule/g. °C = 0.23895 cal₁₅/g. °C; of p = 1 kg*/cm² = 0.9678 atm

Table 121.—Various Isopiestic Thermal Data for Water

 C_p = specific heat at constant pressure = limit approached by the ratio $(\Delta q/\Delta T)_p$ as ΔT approaches zero, Δq being the heat that must be added to the substance in order to increase its temperature by the amount ΔT ;

^{*} These supersede Fig. 41 of his earlier paper 377 which, contrary to these, indicates that at the lower temperatures $(c_p - c_v)$ has a pronounced maximum at a pressure near 5000 kg*/cm².

⁸⁷⁷ Bridgman, P. W., Proc. Amer. Acad. Arts Sci., 47, 439-558 (550) (1912).

Table 121-(Continued)

H= heat content (enthalpy), $H_0=\int_0^T C_p dT$; G=H-ST is the function that Gibbs denoted by ζ and that has been called the "free energy at constant pressure," $G_0=-T\int_0^T (H_0/T^2)dT$; $S_0=$ entropy. All these

quantities refer to the gfw, and those with subscript $_0$ are measured from 0 °K; the pressure is 1 atm; i = ice, w = water. For method employed in extrapolating C_p to 0 °K, see articles cited. The compiler has changed the units, and derived S_0 from H_0 and G_0 . 1 gfw = 18.0154 g; 1 joule/gfw = 0.0551 j/g; 1 cal = 4.185 j.

Unit of C_P and $S_0 = 1$ j/(gfw.°K); of H_0 and $G_0 = 1$ kj/gfw. Temp. = T°K

Ref.a→		Si	mon-			Miet	hing	
T	C_p	H_0	$-G_0$	S_0	′ C _P	H_0	$-G_0$	S_0
273i	41.0	5.35	5.00	37.9	50.2	5.49	5.02	38.4
273vv	75.7	11.36	5.00	59.9	75.4	11.50	5.02	60.5
280	75.4	11.89	5.42	61.8	75.4	12.03	5.45	62.4
290					75.4	12.78	6.09	65.0
300	75.3	13.40	6.55	66.5	75.3	13.53	6.74	67.5
32 0					75.3	15.05	8.15	72.6
340	75.4	16.41	9.40	76.0	75.3	16.56	9.65	77.1
360					75.3	18.07	11.24	81.4
373	75.7	18.91	12.02	92.9	75.3	19.04	12.30	84.0

* References:

Miething, II., Abh. d. Deuts, Bunsen Ges., No. 9 (1920), based upon the data of Nernst, W., Ann. d. Physik (4), 36, 395-439 (1911) and of Pollitzer, F., Z. Elektroch., 19, 513-518 (1913). Simon, F., "Handb. d. Physik" (Geiger and Scheel), vol. 10, p. 363, 1926, based on his own previously unpublished observations.

Table 122.—Enthalpy of Compressed Water

For observations through the critical region, see Table 53.

The enthalpy (H) of a substance is defined by the relation $H = (E + pv) - (E + pv)_0$, where $(E + pv)_0$ is the value of (E + pv) for some state of the substance arbitrarily selected as the basis of reference. For water, the reference state is that of saturation at 0 °C. E is the intrinsic energy, v the volume, and p the pressure; H, E, and v each refers to a unit mass of the substance. The specific heat at constant pressure is $c_p = (dH/dt)_p$.

In some cases the value tabulated is the excess of the corresponding value of H, expressed in Int. steam calories per gram, above the numerical value of the corresponding Centigrade temperature. For example, in Section II, at 75 °C and $p=50 \text{ kg*/cm}^2$ observer S found H to be 75+1.0=76.0 Int. steam cal/g = 318.1 Int. joule/g.

Table 122—(Continued)

I. Pressure = 1 atm. N. S. Osborne. 370

Unit of H either = 1 Int. joule/g or 1 Int. steam cal/g = 4 18605 Int. joules/g, as indicated Temp. = t °C (Int. scale)

Unit→	$_{H}^{\mathrm{Joule}}$	$\operatorname{Cal}_{10^4(H-t)}$	$U_{nit} \rightarrow t$	Joule H	$\operatorname{Cal}_{10^4(H-t)}$
0	0 1026	245	55	230 228	-12
5	21 147	517	60	251 140	54
10	42 126	634	65	272 064	-70
15	63 064	652	7 0	293 000	 55
20	83 976	610	75	313 952	-5
25	104 874	532	80	334 920	+86
30	125 763	435	85	355 907	+223
35	146 650	331	90	376 917	+412
40	167 538	229	95	397 950	+659
45	188 428	134	100	419 011	+971
50	209 324	53			

II. Four sets of values, each indicated by the initial of the experimenter.

	Unit o	f p =	1 kg*	/cm²,	of H	<i>T</i> = 1	Int	steam	cal/g	= 4 1	860	înt j	oule/g	Te	mp i	= °C	:
$t\rightarrow$	•	0	20	25	40	50	60	75	80 - H -	100	120	125	1 40	150	160	175	180
₽ 50	HMª K S TS	1 18 1 2	1 16 1 18	10	1 06 1 02	09	0 96 1 18	1 0	0 89	09	1 1 1 05	11	1 4 1 40	1 5	1 9 2 00	22	2 6 2 82
100	HM K S TS	2 36 2 2	2 9 2 30 2 25	22	2 15 2 0 2	20	1 98 1 88	19	1 86 1 78	1 9 1 8 1 7 1 78	1 9 1 90	20	2 1 2 18	2 3	2 5 2 62	3 0	3 2 3 45
150	HM K S TS	3 53 3 6	3 43 3 38	3 3	3 23 3 12	3 0	3 01 2 92	29	2 82 2 78	2 28 2 6 2 75	2 7 2 75	29	2 8	3 1	3 1 3 38	3 9	3 7 4 00
200	HM K S TS	4 71 4 9	4 55 4 57 4 48	4 3	4 31 4 18	40	4 03 3 95	3 9	3 78 3 72	3 8 3 6 3 5 3 65	3 5 3 62	3 9	3 6 3 78	3 7	3 8 4 09	4 5	4 2 4 60
250	HM K S TS	5 89 6 0	5 70 5 58	5 7	5 40 5 20	5 0	5 06 4 90	4 9	4 74 4 68	4 5 4 3 4 50	4 3 4 50	4 6	4 3 4 62	4 5	4 4 4 85	5 3	4 7 5 25
300	HM K S TS	7 07 7 0	6 79 6 83 6 65	68	6 48 6 25	60	6 09 5 92	5 8	5 70 5 60	5 54 5 4 5 2 5 45	5 1 5 3	5 0	5 0 5 40	5 3	5 0 5 58	5 7	5 3 5 90
350	HM K S TS	8 2		76		69		6 5		6 1		5 7		6 1		61	
400	HM K S TS	8 92 9 2		8 5		8.0		7.3		7 22 7 0		68		68		7 0	

Table 122—(Continued)

$t \rightarrow p$		200	220	225	240	250	260	275	280	H - t	310	320	325	330	340	350	360	370
	HM K S TS	3 9 3 8 4 00	5 6 5 52	60	78	9 2	10 9											·
100	HM K S TS	4 1 4 3 4 3 4 42	5 8 5 78	64		92	10 7	13 8	14 6 15 48	20 7 20 4 21 2 21 85								
150	HM K S TS	4 7 4 8 4 85	6 0 6 08	69	7 9 7 88	90	10 5	13 3	14 1 14 85	19 2 19 6 20 22	22 5	26 7	31 0	32 3	40 1			
200	HM K S TS	5 05 5 1 5 3 5 32	6 3 6 50	70	8 0 8 10	9 4	10 4	13 0	13 7 14 32	18 2 18 2 18 7 19 00	21 1	24 6	28 5	29 0	34 9	42 6 42 9 48	55 6 64	
250	HM K S TS	5 6 5 9 5 98	6 6 6 95	7 5	8 0 8 32	9 6	10 2	12 9	13 2 13 75	17 4 18 0 17 95	20 0	23 1	26 7	26 8	31 4	37 8 37 4 41	46 1 52	58 7 65
	HM K S TS	6 0 5 9 6 5 6 45	6 8 7 42	7 8	8 32 8 72	98	10 1 10 45	12 8	12 8 13 50	16 7 16 7 17 4 17 05	19 0	21 8	25 2	25 0	28 9	33 9 33 7 37 5	40 5 44	48 4 52
350	IIM															31 45		
	K S TS	70		8 0		10 1		12 5		16 9			24 2			35	41	46
400	НМ	69								16 2						29 3		
	K S TS	7 6		8 5		10 3		12 0		16 6			23 1			33	37	42

a References .

Table 123.—Entropy of Compressed Water

For the excess of the entropy of water at 25 °C above that of ice at 0 °K, see Table 207.

The excess of the entropy of water at the indicated temperature (t) and pressure (p) above that of saturated water at 0 °C is S.

Two sets of data are given, distinguished by the initials of the experimenters.

Unit of $p = 1 \text{ kg}^*/\text{cm}^2$, of $S = 1 \text{ millical/g }^\circ\text{K}$ (1nt steam cal) = 4 1860 millipoule/g $^\circ\text{K}$.

			- City					
50	100	150	200	250	300	350	400	Ref a
01					03	02	02	K
70 7	706	70 5	70 4	703	702			K
87 5	87 2	868	86 6	86 2	85.8	85 4	85 2	S
	0 1 0 1 70 7	01 01 01 02 707 706	01 01 02 01 02 02 707 706 705	50 100 150 200 0 1 0 1 0 2 0 3 0 1 0 2 0 2 0 3 70 7 70 6 70 5 70 4	50 100 150 200 S 250 0 1 0 1 0 2 0 3 0 3 0 1 0 2 0 2 0 3 0 3 70 7 70 6 70 5 70 4 70 3	50 100 150 200 5 250 300 01 01 02 03 03 03 03 01 02 02 03 03 04 707 706 705 704 703 702	50 100 150 200 S 250 300 350 01 01 02 03 03 03 03 01 02 02 03 03 04 02 707 706 705 704 703 702	50 100 150 200 S 250 300 350 400 01 01 02 03 03 03 03 04 02 02 707 706 705 704 703 702 702 703 703 703 704 703 704 703 704 703 704 703 704 703 704 703 704 703 704 703 704 703 704 703 704 703 704

HM Havlicek, J., and Miškovsky, L., Helv Phys Acta, 9, 161 207 (Tabelle 1) (1936).
 K W Koch 107
 S Schlegel, F., Z techn Phys
 TS M Trautz and H Steyer 101
 14, 105 107 (1933)

^b Koch's 22 at p = 150, t = 100 is surely wrong; probably it should be either 28 or 27.

			Та	ble 123-	-(Continu	ed)		
<i>p</i> →	50	100	150	200	250 S	300	350	400
40	136.4	136.1	135.9	135.6	135.3	135.0		
50	167.5	166.8	166.3	165.2	165.4	164.7	164.3	163.8
60	197.9	197.5	197.1	196.6	196.2	195.7		
75	241.7	240.9	240.1	239.4	238.7	238.0	237.3	236.6
80	256.1	255.5	254.9	254.3	253.6	253.0		
100	310.6	309.8	308.9	308.1	307.3	306.4		
	310.9	309.9	309.1	308.2	307.4	306.4	305.8	304.9
20	363.9	362.8	361.8	360.8	359.7	358.7		
25	37 6.0	375.0	373.8	372.8	371.8	370.8	369.8	369.7
40	414.2	412.9	414.7	410.4	409.2	407.9		
50	438.1	436.8	435.6	434.3	433.3	431.9	431.0	430.0
60	462.6	461.2	459.7	458.2	456.8	455.3		
75	497.3	495.8	494.2	492.9	491.4	490.0	488.7	487.3
80	509.6	507.9	506.2	504.5	502.8	501.1		
00	555.4	553.4	551.4	549.4	547.5	545.5		
	555.7	552.9	551.0	549.4	547.9	546.2	544.7	543.2
20	600.5	597.9	595.6	593.3	591.0	588.8		
25	610.4	608.1	606.0	604.0	602.0	600.0	598.2	596.7
40	644.4	641.6	639.0	636.3	633.7	631.1		
50	665.5	662.8	660.2	657.7	655.4	652.9	651.0	648.9
60	688.5	685.4	682.2	679.2	676.1	673.1		
75		717.6	714.2	710.9	707.8	704.9	702.2	699.8
30		729.5	725.8	722.1	718.5	714.9		
00		775.1	770.2	765.7	761.4	757.3		
		775.1	770.2	765.7	761.9	757.9	754.6	751.5
10			793.3	788.0	783.2	778.7		
20			817.5	811.0	805.5	800.4		
25			833.9	826.8	821.0	815.5	811.2	806.3
30			843.5	835.1	828.4	822.6		
40			872.9	861.1	852.4	845.4		
50				890.3	878.2	869.2		
-				897.5	885.8	876.1	869.5	863.0

^{*} References:

360

370

Table 124.—Heat of Isothermal Compression of Water

919

954

941.

S

886

907

894

917

903

930

Adapted from a compilation by J. R. Roebuck, 378 based on a graph by P. W. Bridgman. 372, Fig. 7

Q is the amount of heat that must be removed to keep the temperature unchanged when the pressure is increased from 1 atm to 1 atm + p.

K W. Koch.³⁶⁷
 S Schlegel, E., Z. techn. Phys., 14, 105-107 (1933).

⁸⁷⁸ Roebuck, J. R., Int. Crit. Tables, 5, 147 (1929).

Table 124—(Continued)

	Unit of #	= 1 kg	*/cm² =	0.9678	atm; of	$Q_0 = 1 \text{ cal}_1$	g; of	$Q_j = 1$	joulc/g	
$t\rightarrow$	0	20	40	60	80	0	20	40	60	80
Þ			— Qo -					Qs		
500	0.2	0.7	1.5	2.1	2.6	0.8	2.9	6.3	8.8	10.9
1000	0.6	1.6	2.9	4.1	5.0	2.5	6.7	12.1	17.2	21.0
2000	1.9	3.8	5.8	7.9	9.2	8.0	15.9	24.3	33.1	38.5
3000	4.0	6.4	8.7	11.4	13.1	16.8	26.8	36.4	47.8	54.9
4000	6.4	8.9	11.6	14.6	16.5	26.8	37.3	48.6	61.2	69.1
6000	10.6	14.0	17.3	20.9	23.2	44.4	58.7	72.5	87.6	97.2
8000		19.6	23.1	27.0	29.3		82.1	96.8	113.0	122.8
10000			28.7	32.9	35.3			120.2	137.8	147.9
12000			34.5	38.8	40.8			144.6	162.6	171.0

Table 125.—Decrease in Internal Energy of Water on Isothermal Compression

Adapted from P. W. Bridgman. 372, Fig. 8

D= resultant decrease in the internal energy of water when the pressure on it is isothermally increased from 1 atm to $p \text{ kg*/cm}^2$. The work (W) done on the water during such compression is the excess of the heat (Q) given out (Table 124) above D; $W=Q_f-D$.

Un	it of $p = 1$	$kg^*/cm^2 = 0.$.9678 atm;	of $D=1$ jo	ule/g
$\stackrel{t\rightarrow}{p}$	0	20	40	60	80
500 1000 2000 3000	0.2 0.9 3.3 6.5	2.9 5.7 10.5 15.3	5.2 10.0 18.6 25.1	8.4 15.5 27.2 35.6	10.3 18.8 32.2 42.3
4000	9.2	19.0	30.1	42.5	49.8
5000 6000 7000 8000 9000	10.7 10.3	22.0 24.3 26.6 29.7	34.3 38.1 41.5 44.6 47.5	48.1 53.0 57.1 60.9 64.4	56.1 61.1 65.5 69.1 72.4
10000 11000 12000			50.2 52.8 55.0	67.4 70.1 72.4	75.1 77.4 79.5

Joule-Thomson Coefficient for Water.

The Joule-Thomson coefficient (μ) is the decrease in temperature per unit drop in pressure, the expansion being adiabatic. It measures the internal latent heat of expansion, and is the increase in temperature on adiabatic compression.

The several sets of observations given in Table 126 are discordant, and it is to be noticed that at the lower temperatures the observed (dynamic) values (μ_0) differ significantly from the corresponding (static) ones (μ_0) computed from the specific heat and the coefficient of thermal expansion; see discussion in text, p. 256.

The experimental determination of μ has been discussed by K. J. Umpfenbach ⁸⁷⁹ and by G. Tammann. ³⁸⁰

As the temperature is varied, the pressure limits remaining unchanged, a temperature (τ) may be found at which μ passes through zero, changing its sign. This is called the inversion temperature? The following values, τ_1 and τ_2 , were computed by W. Koch ³⁶⁷ and by M. Trautz and H. Steyer, ³⁷¹ respectively, from their determinations of the enthalpy:

þ	50	100	150	200	250	300	kg*/cm²
T1	242.2	244.3		247.3		248.9	°C
72	245.0	249.3	253.8	258.1	263.1	267.3	°C

Table 126.—Joule-Thomson Coefficient for Water

(See text also.)

 $\mu = (dt/dp)_a$, or $(\Delta t/\Delta p)_a$, where Δp is of the order of 100 kg*/cm²; $\mu_o = \mu_o + \delta$, where μ_o is the observed (dynamic) value, and μ_o is the corresponding (static) value computed from the specific heat and the coefficient of thermal expansion; p is the mean of the initial and the final pressure; ()_a indicates that heat is neither added nor removed from the water.

Unit of $p = 1 \text{ kg*/cm}^2$; of μ and $\delta = 0.01 \text{ °C per } 100 \text{ kg*/cm}^2$

I. Tammann and Elbrächter.a

p p	_ 0°°C _	δ	p	_ 30 °C	δ	_ t	70°C	δ
2850	20.8	+6.5	2918	22.2	+11.5	274	5 25.0	+8.6
2712	17.4	+9.3	2734	21.5	+11.7	2500	33.0	+1.1
2572	20.2	+5.9	2554	22.2	+10.7	234		+2.4
2461	19.0	+6.4	2388	21.8	+10.7	2192		-0.1
2351	18.5	+6.6	2244	24.0	+8.2	2072	2 33.6	-1.7
2216	19.2	+5.3	2118	24.4	+7.4	1951	32.3	+4.4
2071	17.9	+5.9	1984	20.9	+10.5	1816		+ 1.6
1942	19.2	+4.0	1806	25.5	+5.4	1676		+3.4
1824	19.5	+2.5	1641	20.6	+9.7	1544		+2.4
1704	18.2	+2.6	1501	20.3	+9.5	1418	3 42.5	-2.2
1580	19.0	+0.7	1350	21.8	+7.4	1290		-0.2
1452	18.0	0.0	1202	23.0	+5.6	1148		-0.6
1325	16.5	+0.1	1022	24.0	+3.8	1020		-1.3
1188	16.6	-0.1	864	22.1	+4.3	886		-0.7
1038	15.2	+0.2	722	23.9	+1.5	752		-2.0
838	13.8	-0.8	560	22.2	+ 3.3	611		-2.1
762	12.8	-0.9	488	24.8	+ 0.1	500		-1.2
630	10.9	-0.3	415	19.6	+5.2	431		-1.7
462	10.2	-2.2	300	23.9	+0.4	294		-2.6
285	8.2	-4.4	210	23.1	+0.6	158		-2.9
98	-0.7	+0.2	125	22.7	+ 0.4	50		-2.4
560	12.8	-2.9	46	22.6	-0.1	886		-2.5A
440	11.9	-4.5				752		-2.4A
330	9.8	-5.0				611		-0.2A
255	8.6	-5.2				506		+3.0A
198	2.9	-2.7				431		+2.3A
88	0.3	-0.9				294		+1.7A
						158		-1.9A
						56	50.3	-0.3A

ero Umpfenbach, K. J., Z. techn. Physik, 12, 25-29 (1931).

⁸⁸⁰ Tammann, G., "Über die Beziehungen zwischen den inneren Kräften und Eigenschaften der Lösungen," Leipzig, Voss, 1907; see Roebuck, J. R., Int. Crit. Tables, 5, 146, 147 (1929).

Table 126-(Continued)

I	I. P	ushin	and G	reben	shchik	$ov.^b$	III. Bridgman. ^c (ICT)					
$t \rightarrow p$	0	25	37 — μο —	54	80		t p	0	20	40	60	80
1 500 1000 1500 2000 2500 3000 3500 4000	-13.0 -2.0 +6.4 11.6 15.0 17.3 18.9	+6.6 +13.0 16.7 18.8 20.3 21.3 22.3 24.2 24.0	+26.0 27.3 27.9 27.9 27.9 27.9 27.9 28.4 29.3	+39.0 37.1 35.7 34.4 33.5 32.9 32.5 32.2	+49.2 46.8 44.5 42.3 40.6 39.2 38.2		1 500 1000 1500 2000 3000 4000 6000 8000 10000 12000	-1.6 +6.8 13.2 18.3 21.5 25.1 26.0 19.4	+13.7 17.5 22.0 24.8 26.3 28.0 28.3 28.9 35.5	28.7 30.0 30.9 31.6 32.2 32.5 32.3 33.6 33.3 33.0 32.0	41.7 41.7 41.3 40.6 39.7 38.1 36.6 34.9 33.7 33.0 32.0	54.8 50.0 46.2 42.7 40.3 36.7 34.4 30.8 27.9 25.7 23.8
I	V. V	V. Ko	ch.867									
	$t \rightarrow p$	0	10	00	200	240 με	250	260	30	00	350	
2	50 00 00 00 00	-235 -235 -236 -237	-12 -12 -13	78 79	-72 -74 -76 -79	-2 -8 -13 -16	+20 +12 +6 +2	+45 +35 +27 +23	+2 +1 +1	38	+702 +399	

- V. By an optical method Mascart 381 found for water at 16 °C and p about 2, $10^4 \mu = 11 \, ^{\circ}\text{C/atm}$.
- P. G. Tait, 382 using a Cu-Fe thermocouple, found the following values (if his "ton" = 2240 lbs) for water at 15.5 °C; he stated that they are to be accepted with caution:

"G. Tammann and A. Elbrächter. 301 The pressure was raised to about 3000 kg*/cm², and then reduced by a series of sudden releases, the decrease in temperature being observed for each step. Each step was about 150 kg*/cm². They believe that the error in μ₀ in no case exceeds ±0.015 °C per 100 kg*/cm² at 0 °C and at 30 °C, nor ±0.030 at 70 °C. In computing μ₀, they used P. W. Bridgman's values for the thermal expansion, and for the lower pressures at 70° Amagat's also; the values from this second computation are here indicated by A.

N. A. Pushin and E. V. Grebenshchikov, 302 included in part in the compilation by J. R. Roebuck (ICT). 304 It will be noticed that these values differ in the same general way as do those of Tammann and Elbrächter from the corresponding ones computed.

way as do those of Tammann and Elbrächter from the corresponding ones computed from Bridgman's data.

^o P. W. Bridgman ^{872, Fig. 14} as given by J. R. Roebuck. ⁸⁸⁴

Table 127.—Heat Liberated by Adiabatic Compression of Water Adapted from M. Trautz and H. Steyer. 371

The heat liberated by adiabatic compression is μc_p , where $\mu = (\delta t/\delta p)_a$ is the Joule-Thomson coefficient, and cp is the specific heat at constant pressure.

²⁸¹ Mascart, Compt. rend., 78, 801-805 (1874).

⁸⁸³ Tait, P. G., Proc. Roy. Soc. Edinburgh, 11, 217-219 (1882). 888 Pushin, N. A., and Grebenshchikov, E. V., J. Chem. Soc. (London), 123, 2717-2725 (1923).

²⁰⁰⁴ Roebuck, J. R., Int. Crit. Tables, 5, 146 (1929).

			-	able in	(00,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	/			
		Unit e	of $p=1$	kg*/cm2; c	$f \mu c_p = 10^{-4}$	joule/g(kg*/cm²)		
$r \rightarrow$	Prat	100	200	300	$p \rightarrow$	Past	100	200	300
8		μ	Ср —		t		/	LCp -	
10	959	950	946	942	120	691	707	720	733
20	923	921	917	913	140	628	649	670	678
30	898	895	893	890	160	548	586	603	628
40	874	871	872	871	180	456	502	532	578
50	851	850	850	850	200	352	398	440	490
60	827	828	830	830	220	222	272	335	398
70	804	808	810	813	240	54	100	184	272
80	781	787	793	797	260	-172	-134	-33	-84
90	758	768	774	782	280		-377	-272	-159
100	736	747	758	768					

Table 127—(Continued)

Table 128.—Isentropic Increase in the Temperature of Water from Saturation to and above the Critical Pressure

Adapted from a table computed by J. H. Keenan.³⁸⁵ S = excess of entropy above that at 0 °C and 1 atm; $p_{\text{crit}} = 218.39$ atm, $t_{\text{crit}} = 374.15$ °C (see Table 241).

Example: For saturated water, S = 0.1 when t = 28.73 °C, the corresponding pressure being 0.037 atm; if the pressure is increased from 0.037 to 218.39 atm, the temperature must at the same time be increased by 0.45 °C (i.e., to 29.18 °C) if S is to remain unchanged.

Unit of $p = 1$	atm; of $S=1$	Int. cal/g.°C = 4.186	joule/g.°C;	of that and	$\Delta t = 1 ^{\circ}C$
$f \rightarrow$	218.39	387.18			
S		Δt	Pant	funt	
0.1	0.45	0.86	0.037	28.73	
0.3	1.39	2.48	0.851	95.59	
0.5	2.76	4.91	9.284	175.28	
0.7	4.94	9.31	50.32	265.19	
0.8	5.68	12.45	96.19	309.08	
0.9	5.64		158.18	347.35	

Table 129.—Specific Heat of Sea-Water

The values in the last pair of columns have been derived by O. Krümmel ³⁸⁶ from those published by J. Thoulet and A. Chevallier ³⁸⁷ and given in the preceding columns.

r= ratio of the specific heat of sea-water to that of pure water at the same temperature; s= salt content; $\rho=$ ratio of the density of sea-water to that of pure water at the same temperature. In all cases the temperature was 17.5 °C.

Unit of $s=1$	g salt per ka	g sca-water; /	and r are ratios	. Tem	p. = 17.5 °C
ρ	7	ρ	*	5	r
1.0025	0.986	1.0275	0.931	0	1.000
1.0050	0.977	1.0300	0.927	5	0.982
1.0075	0.968	1.0325	0.924	10	0.968
1.0100	0.963	1.0350	0.921	15	0.958
1 0125	0.957	1 0375	0.917	20	0.951

⁸⁸⁵ Keenan, J. H., Mech. Eng., 53, 127-131 (1931).

⁸⁸⁶ Krümmel, O., "Handb. d. Ozeanog.," Vol. 1, 1907.

²⁸⁷ Thoulet, J., and Chevallier, A., Compt. rend., 108, 794-796 (1889).

Table	129-	(Continued)
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ø	<i>r</i>	Þ	7	s	7
1.0150	0.952	1.0400	0.913	25	0.945
1.0175	0.948	1.0425	0.910	30	0.939
1.0200	0.944	1.0450	0.907	35	0.932
1.0225	0.940	1.0475	0.903	40	0.926
1.0250	0.935	1.0500	0.900		•

35. THERMAL CONDUCTIVITY OF WATER

At the time that T. Barratt and H. R. Nettleton prepared their compilation,³⁸⁸ there was no available determination of the thermal conductivity of water at temperatures above 100 °C, and they concluded that in the

Table 130.—Thermal Conductivity of Water

(For sea-water, see Table 132.)

The values attributed to B, BN, KH, and ML have been computed by means of their linear formulas as given below; those attributed to SS have been read from their curve. At temperatures below 100 °C the pressure was 1 atm; at higher temperatures it was a few atmospheres greater than the vapor pressure, but the increase produced in the conductivity by the highest pressure used scarcely equals the uncertainty in the observations (cf. Table 131).

Formulas, $\tau = (t - 20)$:

BN° $k = 0.00587 (1 + 0.00281 \tau)$ watt per cm·°C; 0 to 80 °C (ICT) KH° $k = 0.00623 (1 + 0.0012 \tau)$ watt per cm·°C; 0 to 80 °C

 $ML^a k = 0.00610 (1 + 0.0023 \tau)$ watt per cm·°C; 0 to 60 °C $B^a k = 0.00590 (1 + 0.00260 \tau)$ watt per cm·°C; 0 to 80 °C

Unit of $k = 10^{-5}$ watt/cm.° C = 2.389 (10-8)cal/cm.sec.°C. Temp. = t °C.

Ref a→	SS	В	BN	KH	MI.			SS	
t .			k			ŧ	k	*	k
0	554	559	554	608	583	100	680	200	666
10	576	575	570	615	596	110	684	210	659
20	598	590	587	623	610	120	686	220	652
30	615	605	604	630	623	130	687	230	644
40	630	621	620	638	637	140	686	240	635
50	643	636	636	645	650	150	685	250	624
60	654	651	653	653	663	160	682	260	614
70	665	667	670	660	(677)	170	680	270	602
80	671	682	686	668	(690)	180	676	280	590
90	676				, ,	190	672	290	576
100	680					200	666	300	564

⁸⁰⁰ Barratt, T., and Nettleton, H. R., Int. Crit. Tables, 5, 218-233 (218, 227) (1929).

³⁰⁰ Jakob, M., Ann. d. Physik (4), 63, 537-570 (1920).

Schmidt, E., and Sellschopp, W., Forsch. Gebiete Ingenieurw., 3, 277-286 (1932).

Table 130-(Continued)

a References:

- B Bates, O. K., Ind. Eng. Chem., 28, 494-498 (1936). Supersedes Idcm, 25, 431-437 (1933).
- (1933).

 BN Compilation by Barratt, T., and Nettleton, H. R., S88 based upon Bridgman, P. W., Proc. Amer. Acad. Arts Sci., 59, 141-169 (1923); Jakob, M., Sitz. Preus. Akad. Wiss., 1920, 406-413 (1920); Ann. d. Physik (4), 63, 537-570 (1920); Lees, C. H., Phil. Trans. (A), 191, 399-440 (1898); Milner, S. R., and Chattock, A. P., Phil. Mag. (5), 48, 46-64 (1899); Weber, H. F., Sitz. Preus. Akad. Wiss., 1885, 1985, 1985, 185); Repert. d. Physik (Exner), 22, 116-122 (1886), considering Chree, C., Proc. Roy. Soc. (London) (A), 42, 300-302 (1887); 43, 30-48 (1887); Christiansen, C., Ann. d. Physik (Wied.), 14, 23-33 (1881); Graetz, L., Idem, 18, 79-94 (1883); 25, 33-357 (1885); Kohlrausch, F., Diss., Rostock, 1904; Lorberg, H., Ann. d. Physik (Wied.), 14, 291-308 (1881); Mache, H., and Tagger, J., Sitz. Akad. Wiss. Wien (Abt. Ha), 116, 1105-1110 (1907); Wachsmuth, R., Diss., Leipzig, 1892; Ann. d. Physik (Wied.), 48, 158-179 (1893); Weber, H. F., Idem, 10, 103-129, 304-320, 472-500 (1880); Weber, R., Idem (4), 11, 1047-1070 (1903).
- KH Kaye, G. W. C., and Higgins, W. F., Proc. Roy. Soc. (London) (A), 117, 459-470 (1928).
- ML Martin, L. H., and Lang, K. C., Proc. Phys. Soc. (London), 45, 523-529 (1933).
- SS Schmidt, E., and Sellschopp, W. 300

range 0 °C to 80 °C the conductivity can be represented by the formula BN given in Table 130, the accuracy being of the highest and amply sufficient to justify the use of water as a standardizing substance. This formula was based largely on the work of M. Jakob ³⁸⁹ at the Physikalisch-Technischen Reichsanstalt.

Since then, quite different results have been obtained by G. W. C. Kaye and W. F. Higgins at the National Physical Laboratory, and by L. H. Martin and K. C. Lang (see KH and ML, Table 130); and a series extending to 270 °C has been published by E. Schmidt and W. Sellschopp. The last indicates that the variation is not linear in t; that the conductivity reaches a maximum near 130 °C, and has nearly the same value at 300 °C as at 0 °C. These sets of observations were believed to be in error by not more than one or two per cent. Other isolated and less accurate determinations have been published by J. F. D. Smith ³⁹¹ and by T. W. Classen and J. Nelidow. ³⁹²

The theory of the conduction of heat by liquids has been discussed recently by A. Kardos,⁸⁹³ and a series of interesting papers treating of certain thermomechanical properties of liquids and their relations to thermal conductivity has been published by R. Lucas ³⁹⁴ and by F. Perrin and R. Lucas.³⁹⁵

Table 131.—Thermal Conductivity of Compressed Water

Adapted from T. Barratt and H. R. Nettleton ³⁹⁶ and based on P. W. Bridgman. ³⁹⁷

- 891 Smith, J. F. D., Ind. Eng. Chem., 22, 1246-1251 (1930).
- 892 Classen, T. W., and Nelidow, J., Physik. Z. Sowj., 5, 191-199 (1934).
- 888 Kardos, A., Forsch. Gebiete Ingenicurw., 5, 14-24 (1934).
 884 Lucas, R., J. de Phys. (7), 8, 98S-99S, 410-428 (1937); Compt. rend., 204, 418-420, 1631-1632 (1937).
 - sos Perrin, F., and Lucas, R., Compt. rend., 204, 960-961 (1937).
 - 806 Barratt, T., and Nettleton, H. R., Int. Crit. Tables, 5, 218-233 (227) (1929).
 - 897 Bridgman, P. W., Proc. Amer. Acad. Arts Sci., 59, 141-169 (1923).

Table 131—(Continued)

 k_p and k_1 = thermal conductivity of water at the indicated temperature and under the pressures p and 1 atm, respectively; p = 1 atm + P kg*/cm². Temp. = t °C.

	Unit of $P = 1$	kg*/cm2 =	0.9678 atm = 0.9807	megadyne/cm2	
t→ P	$\frac{30}{1000(k_{F})}$	$\frac{75}{(-k_1)/k_1}$	t→ P	$\frac{30}{1000(k_P - $	$75 k_1)/k_1$
1000	58	65	7000	332	345
2000	113	123	8000	366	379
3 000	163	176	9000	398	412
4000	210	225	10000	428	445
5000	253	268	11000	456	476
6000	293	308	12000	Fª	506

a Frozen.

Table 132.—Thermal Conductivity of Sea-water

In 1907, O. Krümmel 398 stated that direct determinations of the thermal conductivity of sea-water were then lacking. He computed a series of values for 17.5 °C and various salinities, based upon the conductivity of pure water and upon the assumption that the heat diffusivity (conductivity divided by product of density times specific heat) is the same for sea-water as for pure water. These values, so corrected as to accord with the value of the conductivity of water given in the *International Critical Tables*, were given in the compilation by T. Barratt and H. R. Nettleton, 399 and are reproduced below. See also J. E. Fjeldstad. 400 Salinity = s grams total salts per kg of sea-water.

	Unit of $k-1$	0-5 watt/cm.°C	; of s -	1 g/kg.	Temp. = 17.5 °C	C
S	0	10	20	30	35	40
k	583	569	563	560	558	557

36. TEMPERATURE OF MAXIMUM DENSITY OF WATER

That the density of water under a pressure of 1 atm is a maximum at a temperature (t_m) near 4 °C has long been known, but the exact value of t_m , 3.98 °C on the international hydrogen scale (3.98 to 4.01 for mercury-in-glass thermometers), was not established until around the beginning of this century.

A list of 36 early and widely varying estimates of t_m has been published by F. Rossetti.⁴⁰¹ The only ones in that period that need be considered are those of Despretz, included in the following table.

As the pressure is increased, t_m decreases, its rate of decrease, at least for the first few hundred atmospheres, being essentially constant and greater than that of the depression of the freezing point. The first attempt to

⁸⁰⁸ Krümmel, O., "Handbuch der Ozeanographie," Vol. 1, p. 280, Stuttgart, J. Engelhorn, 1907.

Barratt, T., and Nettleton, H. R., Int. Crit. Tables, 5, 218-233 (229) (1929).

⁴⁰⁰ Fjeldstad, J. E., Geofysiske Publ., 10 No. 7, 1933.

⁴⁰¹ Rossetti, F., Ann. dc chim. ct phys. (4), 10, 461-473 (1867) ← Atti. Ist. Veneto, 12, (1866).

estimate the variation of t_m with the pressure seems to have been that of I. D. van der Waals. 402 His estimates, based upon an equation which he fitted to 3 values of the compressibility as determined by C. Grassi 311 and upon the thermal expansion as given by the formula of H. Kopp 403 $[10^6(v-v_0)/v_0=-61.045t+7.7183t^2-0.03734t^3, 0^{\circ} \le t \le 25 {\circ} C]$ are quoted in a much used handbook, and ascribed to Grassi, computation by van der Waals. Actually, Grassi is in no way responsible for those values, which, indeed, are discordant with all direct determinations. from which they were derived are unsuited to that purpose, and the equation which van der Waals used for the compressibility reproduces only those three of Grassi's determinations which were used in deriving it, giving at other temperatures values which are entirely impossible. For these reasons, those estimates by van der Waals are not included in this compilation, it being sufficient to remark that for the range 1 to 10.5 atm they lead to a mean depression of t_m of 0.072 °C per atmosphere, nearly three times that found experimentally for a wide range of higher pressures (see Table 133).

The presence of a solute likewise depresses the value of t_m ; again, by more than it depresses the freezing point. To a first approximation, each depression is proportional to the concentration of the solution.

A. J. Bijl ⁴⁰⁴ has reported values of t_m for mixtures of water and finely divided sugar-charcoal. They fall below the value for water, the depression (Dt_m) of t_m depending upon the amount of charcoal; e.g., for 7.87 g charcoal and 15.03 g water, $Dt_m = 4.8 \,^{\circ}\text{C} = 0.61 \,^{\circ}\text{C/g-charcoal}$; for 4.55 g charcoal and 16.82 g water, $Dt_m = 2.7 \,^{\circ}\text{C} = 0.59 \,^{\circ}\text{C/g-charcoal}$. He suggests that the effect arises from the composition of the water (relative amounts of the several polymers) in the absorbed layer differing from that of water in bulk.

Table 133.—Temperature of Maximum Density of Water

(For sea-water, see Table 134.)

If P does not exceed a few hundred atmospheres, the temperature of maximum density (t_m) is approximately given by $t_m = 3.98_2 - a(P-1)$ °C. The earlier observations gave 3.98 to 4.01 when P=1, depending upon the nature of the glass of the thermometer. Lussana a derives from his observations a=2.25 °C per 100 atm, which probably is as good as we can do; that value is higher than the mean of those given below because they are based on $t_m=4.00$ °C for P=1, while his equation calls for 4.10 °C. If P is not too great, the freezing point is $t_f=-0.0075(P-1)$; hence $t_m=t_f$ when P is about 270 atm.

⁴⁰² van der Waals, J. D., Beibl. zu Ann. d. Physik, 1, 511-513 (1877) ← Med. Kon. Acad. Wet. Amsterdam, Afd. Nat. (2), 11, 1-13 (1877).

⁴⁰⁸ Kopp, H., Ann. d. Physik (Pogg.), 72, 1-62, 223-293 (44) (1847).

⁴⁰⁴ Bijl, A. J., Rec. trav. chim. Pays-Bas, 46, 763-769 (1927).

Table 133—(Continued)

Unit of $P = 1$ atm; of $a = 1$ °C per atm. Temp. = t_m °C	Unit	of	P	=	1	atm;	of	a	=	1	°C	per	atm.	Temp.	=	t-m	۰,	•
--	------	----	---	---	---	------	----	---	---	---	----	-----	------	-------	---	-----	----	---

P =	1 atm —				D.G	- £			
tm	Ref.ª	P	t m	100a	Ref.	of pressure	t _m	100a	Ref.
3.982	Best	-26.3	4.6	2.2	M11	166	0.40	2.18	L95
3.982	de C94	-20.5	4.5	2.3	M11	200	-0.44	2.23	L95
3.98	Ch97 ^b	-12.9	4.3	2.2	M11	222	-0.91	2.22	L95
3.98	TSD00	+41.6	3.3	1.7	A93	251	-1.54	2.22	L95
3.98_{a}	de C03	93.3	2.0	2.2	A93	268	-1.82	2.18	L95
3.99	D37	145	0.6	2.4	A93	300	-2.57	2.20	L95
4.00	D39	197	ca.0	2.0	A93	322	-3.05	2.20	L95
4.07	R66	47	3.06	2.0	L95	150-600		2.4	T82
4.04	R68	58	2.75	2.19	L95	2	•	2.0	T88
4.10	W78	100	1.90	2.12	L95	600	0	0.67	PG23
3.96	S92	112	1.68	2.09	L95				
4.05	L95	148	0.77	2.20	L95				
3.972	Ma91	163	0.40	2.22	L95				

a References:

A93	Amagat, E. II., Compt. rend., 116, 946-952 (1893).
de C94	de Coppet, L. C., Ann. de chim. et phys. (7), 3, 240-269 (1894).
de CO3	de Coppet, L. C., Idem. 28, 145-213 (1903).
Ch97	Chappuis, P., Ann. d. Physik (Wied.), 63, 202-208 (1897).
D37	Daniel C. Cart. a. 1 hysik (Wille), 03, 202-200 (1897).
137	Despretz, C., Compt. rend., 4, 124-130 (1837) \rightarrow Ann. d. Physik (Pogg.), 41, 58-71 (1837).
D39	Despretz, C., Ann. de chim. et phys. (2), 70, 5-81 (1839).
L95	Lussana, S., Nuovo Cim. (4), 2, 233-252 (1895).
M11	Meyer, J., Abh. d. Deutsch. Bunsen-Ges., 3, No. 1, whole No. 6, 1911.
Ma91	Makaroff, C. O., J. Russ. Phys. Chem. Soc. (Chem.), 23 II, 30-88 (1891).
PG23	Pushin, N. A., and Grebenshchikov, E. V. 3883
R66	Rossetti, F., Ann. de chim. et phys. (4), 10, 461-473 (1867) - Atti Reg. Ist.
	l'eneto (3), 12, (1866).
R68	Rosetti, F., Atti Reg. Ist. Veneto (3), 13, 1047-1093, 1419-1457 (1868) → Ann. de
	chim. et phys. (4), 17, 370-384 (1869). Each (R66, R68) abstracted in Ann d.
	Physik (Pagg.) Erg. Bd., 5, 258-275 (1871).
S92	Scheel, K., Ann. d. Physik (Wied.), 47, 440-465 (1892).
T82	Tait, P. G., Proc. Roy. Soc. Edinburgh, 11, 813-815 (1882), from observations
	of Marshall, D. H., Smith, C. M., and Omond, R. T., Idem, 11, 809-813 (1882).
T88	Tait, P. G., Beibl. zu Ann. d. Physik, 13, 442-445 (1889) ← "Report Sci. Res.
200	Voy H M S Challenger, Phys. and Chem." 2. Part 4. London, 1888.
TSD00	Thiesen, M., Scheel, K., and Diesselhorst, H., Wiss. Abh. PhysikTechn. Reich-
131700	sanstalt, 3, 1-70 (1900) → Z. Instk., 20, 345-357 (1900).
33770	Weber, L., Beibl. zu Ann. d. Physik, 2, 696-699 (1878) ← Jahresber. Comm.
W78	H iss. Untersch. Deuts. Mecre in Kiel, 4-6 (1874-76), 1-22 (1878).
	11 188. (Intersent Dents, Meere in Aut, 4-0 (10/4-/0), 1-22 (10/6).

⁶ The equations by means of which Chappuis represents his two ultimate series of observations between 0° and 10 °C, lead to $t_m = 3.978$ and 3.994, respectively; and their mean, which defines his definitive values (Table 99), gives $t_m = 3.986$ °C.

Table 134.—Temperature of Maximum Density of Sea-water

D. H. Marshall, C. M. Smith, and R. T. Omond ⁴⁰⁸ have reported that when sea-water (not more particularly specified) is adiabatically expanded from P to 1 atm there is no resultant change in temperature if the associated temperature and pressure (P) have the following values: $-5\,^{\circ}$ C, 153 atm; $-8.5\,^{\circ}$ C, 306 atm; $-11\,^{\circ}$ C, 458 atm; and $-13\,^{\circ}$ C, 610 atm.

The following data all refer to a pressure of 1 atm. It seems probable that all values of t_m in Section I should be increased by about 0.035 °C (cf. value for s = 0 with Table 133); it is believed that the data in that section are in other respects to be preferred to those in Section II.

⁴⁰⁵ Chappuis, P., Trav. et Mem. Bur. Int. Poids et Mes., 13, D1-D40 (1907).

⁴⁰⁰ Marshall, D. H., Smith, C. M., and Ormond, R. T., Proc. Roy. Soc. Edinburgh, 11, 809-813 (1882).

Table 134—(Continued)

In Section II, the values given for the salinity (s) have been estimated from the density by means of Table 109 except as the contrary is indicated, and are, together with Dt_m/s , only approximately correct. It will be noticed that the values (B) derived from the compilation by J. A. Beattie ⁴⁰⁷ do not entirely accord with the others; although they are said to have been based upon the observations by L, Ma, N, R, and W. It seems likely that the relation he used to connect ρ_0 with s differs from that used by the present compiler.

 $\rho_0 \equiv 1 + 10^{-4} \Delta_0$ is the density at $0 \, ^{\circ}$ C; $\rho_{max} \equiv (1 + 10^{-4} \Delta_m)$ is the density at t_m ; $t_m \, ^{\circ}$ C = temperature of maximum density; $Dt_m \equiv (3.947 - t_m)$ in Section I, and $(3.98 - t_m)$ in Section II.

Unit of s=1 g salt per kg sea-water; of ρ_0 and $\rho_m=1$ g/cm³; of $Dt_m/s=0.1$ °C per (g/kg)

I.	O. Krün	mel. ⁴⁰⁸						
s	Δ_m	t_m	Dt_m/s	s	Δ_m	t m		1)tm/s
0 1 2 3 4	0 8.5 16.9 25.1 33.3	3 947 3.743 3.546 3.347 3.133	2.04 2.00 2.00 2.04	20 21 22 23 24	160 7 168.7 176.7 184.8 192.9	-0.310 -0.529 -0.744 -0.964 -1.180		2.13 2.13 2.13 2.14 2.14
5 6 7 8	41.5 49.6 57.7 65.8	2.926 2.713 2.501 2.292	2.04 2.04 2.06 2.07 2.07 2.07	25 26 27 28 29	201.0 209.1 217.2 225.3	-1.398 -1.613 -1.831 -2 048		2 14 2.14 2.14 2.14
10 11 12 13 14	73.8 81.8 89.7 97.6 105.6 113.5	2.075 1.860 1.645 1.426 1.210 0.994	2.07 2.09 2.09 2.10 2.11 2.11	30 31 32 33 34	233.4 241.5 249.7 257.8 265.9 274.0	-2.262 -2.473 -2.687 -2.900 -3.109 -3.318		2.14 2.14 2.14 2.14 2.14 2.14
15 16 17 18 19	121.3 129.2 136.9 144.8 152.7	0.772 0.562 0.342 0.124 0.090	2.11 2.12 2.12 2.12 2.12 2.13	35 36 37 38 39	282 2 290.4 298.6 306.8 315.0	-3.516 -3.524 -3.733 -3.936 -4.138 -4.340		2.14 2.13 2.13 2.13 2.13
20 21	160.7 168.7	-0.310 -0.529	2.13 2.13	40 41	323.2 331.4	-4.541 -4.738		2.12 2.12
II.	Various	observers	s.					
s	Δ_0		m/s Ref.a	s	Δ_0	t_m	Dtm/s	Ref.a
9.1 ^b 18.1 ^b 27.1 ^b 36.1 ^b 8	71 139	$ \begin{array}{ccc} 0.5 & 1 \\ -1.3 & 1 \\ -3.2 & 2 \\ +2.2 & 2 \end{array} $.0 B 2 L	42 47 35 35 33 34	335 381 281 281 267	-4.6 -5.3 -4.74 -3.90 -3.21	2.0 2.0 2.5 2.3 2.2	L L N R
1/	139	-0.4 2	.6 L	34	273	-3.67	2.2	D

2.0

2.0

208

W

W

M

⁴⁰⁷ Beattie, J. A., Int. Crit. Tables, 3, 108 (1928).

⁴⁰⁸ Krümmel, O., "Handb. d. Ozeanog.," Vol. 1, 1907.

Table 134—(Continued)

a References:

Beattie, J. A. 407 (based on L, Ma, N, R, and W). Despretz, C., Compt. rend., 4, 435-440 (1837); Ann. d. Physik (Pogg.), 41, 58-71 (1837); Ann. de chim et phys. (2), 70, 5-81 (1839). Lenz, R., Mém. Acad. Sci. Russie (7), 29, No. 4 (1881) (observations by Reszow). Makaroff, C. O., J. Russ. Phys. Chem. Soc. (Chem.), 23 II, 30-88 (1891). v. Neumann, C., Ann. d. Physik (Pogg.), 113, 382 (1861) \leftarrow Diss., München, 1861. Rossetti, F., Atti Reg. Ist. Veneto Sci., Let., ed Arti (3), 13, 1047-1093, 1419-1457 (1868) \rightarrow Ann. dc. chim. et phys. (4), 17, 370-384 (1869) \rightarrow Ann. d. Physik (Pogg.) Erg. Bd., 5, 258-275 (1871). Weber, L., Jahresber. Comm. Wiss. Unters. Deuts. Meere in Kiel, 4-6, (1874-1876), 1-22 (1878) = Diss., Kiel, 1877 \rightarrow Beibl. Ann. d. Phys. (Wied.), 2, 696-699 (1878).

^b Given in the citation.

REFRACTIVITY OF WATER

The data for the refraction of water given in Table 135 are believed to be the best of the kind now available. The reduction of the observations on which they are based was not completed until after the rest of this section had been written; and since they have been received, the remainder of the section has been only slightly revised, mainly by increasing the number of entries for the visible spectrum in Table 137, which initially included only the better values.

Relative to the discussion of a possible dependence of the properties of water upon its recent thermal history (see p. 170+), V. K. LaMer and M. L. Miller 400 have sought for a difference between the index of refraction of water newly boiled, rapidly chilled, and measured at once, and that of water treated in a similar manner but kept at room temperature for three days before measuring the index. No difference was found; the precision was ±3 in 106. It should, however, be remembered that the index of refraction of water exhibits no anomaly at 4 °C,410,411 and that both B. C. Damien 411 and V. S. M. v.d. Willigen 412 have reported that the indices of different samples of water, nominally identical, may differ appreciably. The former stated that, like the latter, he had observed that apparently identical specimens of water in which no impurity could be found chemically may have different indices, although any one given sample always had the same index.

Intercomparison of data obtained by various observers is aided by comparing those of each with the same interpolation formula. C. Chéneveau 413 accepted formula (1), which is due to F. F. Martens,414 as valid at 18 °C and for wave-lengths (λ) in the range $\lambda = 0.224$ to 1.256 μ ; the unit of λ in the formula is 1μ , and the index is with reference to air at atmospheric pressure and at the same temperature as the water.

$$n^2 = 1.76148 - 0.013414\lambda^2 + 0.0065438/(\lambda^2 - 0.0132526)$$
 (1)

410 Jamin, J., Compt. rend., 43, 1191-1194 (1856).

⁴⁰⁰ LaMer, V. K., and Miller, M. L., Phys. Rev. (2), 43, 207-208 (1933).

an Damien, B. C., Ann. Sci. Ecole Norm. Sup. (2), 10, 233-304 (272-278) (1881) $\rightarrow J$. de Phys. (1), 10, 198-202 (1881). 412 v. d. Willigen, V. S. M., Arch. Mus. Teyler, 1, 74-116, 161-200, 232-238 (1868).

For the same unit of λ and under the same conditions except that the temperature is 20 °C, J. Duclaux and P. Jeantet ⁴¹⁵ have given formula (2).*

$$n^2 = 1.76253 - 0.0133998\lambda^2 + 0.00630957/(\lambda^2 - 0.0158800)$$
 (2)

Neither of these formulas fits the observed values satisfactorily if $\lambda < 0.25 \,\mu$; but the second fits the more closely, and its fit can be improved by adding to it the term 10^o where $c = 107.73(0.064156 - \lambda^2) - 5$. That formula (2), as so modified, is used as the norm with which to compare the observed values (Tables 137 and 138). The three formulas are compared in Table 139, where the values of $dn_o/d\lambda$ for each of a number of values of λ are also given. It will be noticed that the dispersion of water in the ultraviolet exceeds that of quartz.

Variation with the Temperature (Tables 140, 141, 142, and 143).

Each of the several measurements of the variation of the index of refraction with the temperature has usually been summarized by a formula. A number of these formulas for water are given and compared in Table 142. Except that of Ketteler, they are all algebraic expressions involving only integral powers of t. Ketteler's is this: $(n^2-1)\cdot(v^*-\beta)=C(1+\alpha e^{-kt})$, where v^* is the specific volume of the water, C is the value of $(n^2-1)v^*$ when the substance is in the gas phase and greatly expanded, and α , β , and k are constants fixed by the substance alone; C varies with λ , determining the dispersion.

It will be noticed that Ketteler's formula can be put in the form $(n^2-1)=f(\lambda)\cdot F(t)$, where f and F are each a function of a single variable, λ or t. Whence, for a given λ , $(n_1^2-1)/(n_2^2-1)=F(t_1)/F(t_2)$ and $(n_1^2-n_2^2)/(n_1^2-1)=1-F(t_2)/F(t_1)$ are each independent of λ . Likewise, for a given t, similar ratios are independent of t. Flatow's extended series of observations on water ($\lambda=2145$ to 5893A, t=0 to 80 °C) does not satisfy these conditions.

Although the temperature coefficient of n is a function of λ , its variation with λ is not very rapid (see Table 141).

There is no evidence of an anomaly at $4 \,^{\circ}$ C; the index continues to increase as the temperature is reduced below that temperature, and B. C. Damien, whose observations extended to $-8\,^{\circ}$ C, found no maximum. Others have, however, found that the index does pass through a maximum at a temperature (t_{mr}) not far from $0\,^{\circ}$ C, but the several observers do not agree regarding that temperature. J. Jamin 416 concluded that t_{mr} was

^{*}A typographic error occurs in the paper cited, the value of the numerator of the third term being given as antilog $\overline{3.00800}$, whereas the computed values for n show that it should have been antilog $\overline{3.80000}$.

⁴¹⁸ Chéneveau, C., Int. Crit. Tables, 7, 13 (1930); Recueil de const. phys. (Soc. fr. de Phys.), Paris, 1913.

⁴¹⁴ Martens, F. F., Ann. d. Physik (4), 6, 603-640 (1901).

⁴¹⁵ Duclaux, J., and Jeantet, P., J. de Phys. (6), 5, 92-94 (1924).

near 0 °C; L. Lorenz ⁴¹⁷ that it was +0.01 °C for the D-line ($\lambda = 5893$ A) and +0.17° for Li ($\lambda = 6708$ A); C. Pulfrich ⁴¹⁸ placed it between -1 and -2 °C, his observations extending to -10 °C; E. Ketteler ⁴¹⁰ placed it at -1.5 °C; and N. Gregg-Wilson and R. Wright ⁴²⁰ at -0.5 °C. L. W.

Table 135.—Index of Refraction of Water in the Visible Spectrum: Preferred Values

(L. W. Tilton and J. K. Taylor. Numerical data privately communicated by Tilton prior to the publication of the detailed account of their work, which contains extensive tables covering the ranges 0 to 60 °C and 4000 to 7250A.)

The index is with respect to dry air at a pressure of 760 mm-Hg and at the same temperature as the water. Under δ are given the values of $10^5(n_o-n)$, n_o being the value defined by the expression given in the head-matter of Table 137, and n being the index here given for 20 °C.

Unit of
$$\lambda = 1 \ \mu = 10^{-4} \ \text{cm} = 10^{4} \ \text{A}$$
. Temp. = $t \ ^{\circ}\text{C}$

I. Index with respect to dry air at the same temperature (t °C) and a pressure of 760 mm-Hg.

$t \rightarrow \lambda$	10	20	n30	40	20 8	15	25 (106) dn/	dt
0.70652 0.66781 0.65628	1.330704 1567 1843	1.330019 0876 1151 2988	1.328993 9843 1.330116 1940	1.327685 8528 8798 1.330610	$ \begin{array}{r} -2.0 \\ -0.2 \\ +0.2 \\ +2.2 \end{array} $	68.8 69.4 69.6 70.7	102.9 103.6 103.8 105.0	130.9 131.7 131.9 133.2
0.58926 0.58756	3690 3744	3041	1993	0662	+2.2 + 2.4	70.7	105.0	133.2
0.57696 0.54607 0.50157	4085 5176 7070	3380 4466 6353	2331 3411 5289	0998 2071 3939	+2.6 +3.3	70.9 71.4 72.1	105.2 105.8 106.7	133.5 134.1 135.2
0.48613 0.47131	7842 8653	7123 7931	6055 6860	4702 5504	+3.3	72.3 72.6	107.0 107.3	135.4 135.8
0.44715 0.43 5 83 0.40466	1.340149 0938 3476	9423 1.340210 2742	8347 9131 1.341656	6984 7765 1.340280	+2.9 +2.6 +1.9	73.0 73.2 73.8	107.8 108.1 108.9	136.5 136.8 137.8

II. Index for the D-lines of Na (Hartmann's mean $\lambda = 0.58926 \,\mu$), with respect to dry air at the same temperature (t °C) and a pressure of 760 nm-Hg.

ŧ	n	ŧ	**	t	22	t	71	t	72	ŧ	**
10	1.333690	15	1.333387	20	1.332988	25	1.332503	30	1.331940	35	1.331308
11	638	16	315	21	897	26	396	31	819	36	173
12	582	17	238	22	803	27	287	32	695	37	036
13	521	18	158	23	706	28	174	33	569	38	1.330896
14	456	19	075	24	60 6	29	059	34	440	39	754

Tilton and J. K. Taylor 421 find for the index with respect to air at the same temperature $t_{mr} = +0.19$ °C for the D-line and +0.33 °C for

⁴¹⁰ Jamin, J., Compt. rend., 43, 1191-1194 (1856).

⁴¹⁷ Lorenz, L., Ann. d. Physik (Wied.), 11, 70-103 (1880).

He($\lambda = 6678$ A), and they compute for the absolute index at these wavelengths the values $t_{mr} = -0.05$ and +0.09 °C, respectively.

Both Jamin and Damien have reported that the act of freezing is preceded by an anticipatory decrease in the index. Damien 411 has described

Table 136.—Reduction of the Index of Refraction of Water from Air to Vacuum

If $n_{\rm air}$ is the index of refraction of water with reference to air at the same temperature and a pressure of 760 mm-Hg, then the index with reference to a vacuum is $n_{\rm vac} = n_{\rm air} + \Delta$, Δ depending upon the wave-length (λ) and the temperature. The following values of Δ are based upon the values found by W. F. Meggers and C. G. Peters ⁴³⁰ for the index of refraction of air; for air at a fixed pressure, (n-1) is inversely proportional to the absolute temperature.

Unit of $\lambda = 1 \mu = 10^4 \text{ A} = 10^{-4} \text{ cm}$. Temp. = t °C Temperature = 20 °C.

1. 16	emperature	$= 20^{\circ}$						
λ	10 ^ε Δ	λ	$10^5 \Delta$	λ	$10^5 \Delta$	λ	105 A	
0.20	45.5	0.24	41.4	0.35	37.8	0.55	36.3	
0.21	44.1	0.25	40.8	0.40	37.2	0.60	36.2	
0.22	43.0	0.27	39.8	0.45	36.8	0.70	36.0	
0.23	42.1	0.30	38.8	0.50	36.5	0.90	35.7	
II. T	he D-lines	$\lambda = 0.5$	5893.					
t	10 ⁵ Δ	t	$10^5 \Delta$	t	10 ⁸ △	1	10 ⁸ Δ	
-1	0 40.4	10	37.5	30	35.0	70	30.8	
	5 39.7	15	36.9	40	33.9	80	29.8	
	0 38.9	20	36.2	50	32.8	90	29.0	
+ .	5 38.2	25	35.6	60	31.7	100	28.1	

the phenomenon thus: "J'ai eu bien souvent l'occasion ... d'observer une brusque diminution de l'indice sans cause apparente. L'image d'une raie étant superposée au réticule, on voyait tout à coup cette image se déplacer lentement et graduellement. Un instant après seulement, des aiguilles de glace se formaient dans le prisme. Comme le fait remarquer M. Jamin: La congélation se prépare pour ainsi dire à l'avance au moment où elle va s'opérer."

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418 Pulfrich, C., Idem, 34, 326-340 (1888).
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⁴¹⁹ Ketteler, E., Idem, 33, 353-381, 506-534 (1888).

⁴²⁰ Gregg-Wilson, N., and Wright, R., J. Phys'l Chem., 35, 3011-3014 (1931).

⁴²¹ Tilton, L. W., and Taylor, J. K., private communication, 1935.

⁴²² Bramley, A., Phys. Rev. (2), 33, 279, 640 (1929); J. Opt. Soc. Amer., 21, 148 (1931).

⁴²³ Chéneveau, C., Ann. de chim. et phys. (8), 12, 145-228, 289-293 (1907).

⁴²⁴ Tilton, L. W., Bur. Stand. J. Res., 2, 909-930 (RP64) (1929); 6, 59-76 (RP262) (1931); 11, 25-58 (RP575) (1933); 13, 111-124 (RP695) (1934); 14, 393-418 (RP776) (1935).

⁴²⁵ Dufet, H., "Recueil de données numériques Optique." Publ. by Soc. Fr. de Physique, Gauthier-Villars, Paris, 1898.

⁴⁹⁶ Korff, S. A., and Breit, G., Rev. Mod. Phys., 4, 471-503 (1932).

⁴²⁷ Tilton, L. W., J. Res. Nat. Bur. Stand., 17, 639-650 (RP934) (1936).

⁴²⁸ Tilton, L. W., and Taylor, J. K., Idem, 18, 205-214 (RP971) (1936).

Tilton, L. W., and Taylor, J. R., J. Res. Nat. Bur. Stand., 20, 419-477 (RP1085) (1938).
 Meggers, W. F., and Peters, C. G., Bull. Bur. Stand., 14, 697-740 (S327) (1918).

Effect of Electric Field. (For Kerr effect, see Section 51.)

When an electric field of high frequency is applied to water in a direction perpendicular to that of the propagation of light through it, certain effects are observed which were initially interpreted as indicating that water has a set of absorption bands (and consequently, exhibits anomalous dispersion) for waves 3 to 10 meters long. Later observations showed that such an explanation is incorrect.⁴²²

Table 137.—Various Values of the Refraction of Water at 20 °C

(For the preferred values in the Visible Spectrum, see Table 135; for reduction to vacuum, see Table 136; for a comparison of certain sets of values, see Table 138; for $\lambda > 5000 \,\mu$, see Table 172.)

These indices (n) are with reference to air at a pressure of approximately one atmosphere and at the same temperature as the water, usually 20 °C, but those quoted from Rubens (Rub) are for 12 °C, and those from Rubens and Ladenburg (RL) are for 18 °C. In these exceptional cases the precision of measurement is not great enough to justify a correction for so small an interval as 8 °C. If several sources are cited for the same value, that value is the mean of those from the several sources, and only rarely does any one of the individual values differ from that mean by so much as 2 in the fifth decimal place. If the group of references includes ICT, then the tabulated mean is exactly that given by Chéneveau in the International Critical Tables. For certain frequently studied wave-lengths, several values of n are given, the first being regarded as superior to the others. No distinction has been made between the values of n that have been determined absolutely and those that have been derived from such absolute values by means of relative measurements, either by the same or by another observer.

The somewhat arbitrarily chosen norm with which the several values are compared is

$$(n_o)^2 = 1.762530 - 0.0133998 \lambda^2 + \frac{0.00630957}{\lambda^2 - 0.0158800} + \frac{10^{[107.731(0.084156 - \lambda^2) - 5]}}{10^{[107.731(0.084156 - \lambda^2) - 5]}}$$

the unit of λ is 1μ (see text). The values of λ used in computing n_c are those here given, those given to 5 significant figures having been taken from the list of wave-lengths given by H. Kayser.⁴³¹

It will be noticed that the ICT values agree with the DJ ones until $\lambda = 0.214 \,\mu$ is reached, where a sudden break of 40 in the fifth decimal place occurs; with increasing values of λ , the discrepancy decreases until at $\lambda = 0.397 \,\mu$ the two series again coincide, and thereafter continue to coincide to the end of the DJ series.

⁴⁸¹ Kayser, H., Int. Crit. Tables, 5, 276-322 (1929).

Table 137-(Continued)

Unit of $\lambda = 1 \mu = 10^{-4} \text{ cm} = 10^4 \text{ A}$

			Uı	nit of $\lambda = 1$	$\mu = 10^{-6}$	cm = 10	A	
	— х ——	- "	105(no-	n) Ref.a		λ	n	108 (nc-n) Ref. a
Cub		0.99999652	(Th	Fe	83		+16.5 Du
Cu	0.00015	4 0.99999034			CI	0.4415	7 981	+ 2.0 ICT, F1
Cu	0.00013	4 0.9999964		St	Çq	0.4415	945	
Cu	0.00013	4 0.9999963		Sm	He	715		
A	0.1151	Mete		ICT, DJ	Cd	0.46783		+ 1.4 ICT, F1
Ag	0.1829	1.46379	-41.9	ICT, DJ	Ca	78.	817	- 0.6 Si
Ag	0.1832	264	-18.2	Dî	Çq	0.47999		- 1.9 ICT, FI
Ag	0.1834	199	-19.8		Cd	999		
Ag	0.1835	141	+ 7.5		Нβ	0.4861.	714	+ 1.6 ICT, OL, Sch, Si + 0.6 Wil, Wil
Ag	0.1838	1.46060	- 3.6					Sch, Si
Ag	0.1839	013	+12.5		Нβ	613	715	+ 0.6 Wil, Wil
Ag	0.1849	1.45715	+11.8	DĴ	Hβ	0.48613		- 3.4 Br
Ag	0.1853	595	+16.3	DJ	118	613		-22.4 Ka
Al	0.18547	528	+34.3		IJβ	613		+ 5.6 La
Al	0.18582	477	-13.9	DI	Нβ	613	704	+11.6 Be, Da,
A1	627	343	- 5.0	ICT, DJ				Du
Αl	0.19352	1.43595	+ 4.4	DJ ICT, DJ	N	0.50032		- 0.3 DJ - 6.8 Si
Al	898	1.42572	5.9	ICT, DI	Cd N	858		- 6.8 Si
Zn	0.20255	1.41993	- 7.0	1)1	N	0.51795	567	- 3.0 DJ
Zn	619	1459	- 2.1	DĴ	Mg	0.51836	549	+13.2 Du
Zn	0.21385	1.40500	+ 7.9	DĴ	Cd	0.53380		- 1.3 ICT, F1
Cd	444	437	+ 5.6	DĴ	Tl	505	490	+ 2.6 1CT
Cd	444	397	+45.9	IČT, FI	Tl	505		+ 0.6 Br, Sch
Al	740	128	+ 0.8	DI.	TI	505	498	- 5.4 Si
Cd	946	1.39883	+40.4	DJ ICT, Fl	TI	505	485	+ 7.6 Rü
Al	0.22100	775	+ 1.5	DJ T	Ti	505	481	+11.6 Du, Ket,
Al	636	305	+ 0.4	DĬ				Wm
Cd	650	257	+34.2	ICT, FI	Hg	0.54607	447	4 2.9 ICT
Al	0.22691	1.39258		DI.	Hg	607	4485	+ 1.4 Ja
Ĉď	0.23129	1.38878	-0.5 +35.6	DJ	Hg	607	440	+ 9.9 Ro
ΑÏ	671	533	- 5.2	ĨĊT, FI	N	0.56795	370	+ 1.0 DJ
Hg	783	233	1 10 4	Dl	Hg	0.57696	342	- 1.4 IČT
Au	0.24280	434	+19.4	Ro	Hg	696	3402	+ 0.4 Ja
		103	+ 37.1	ICT, FI	Hg	907	333	+ 0.8 ICT
Hg	827	1.37809 1.37734	+17.5	Ro	Hg	907	333 ₅	+ 0.3 la
Zn	0.25020		-11.2	DJ	He	0.58756	305	+ 1.5 Ro
Al	680	406	-13.2	DJ	Na	0.58929	300	+ 1.0 ICT, F1,
Çd	730	349	+20.0	DÍ ICT, FI			•••	Sch
Al	753	372	-12.6	173	Na	0.58929	1.33299	+ 2.0 BBD, HP,
Hg	0.2576	338	+17.3	Ro		0.00,00	1.00277	Ve, Wa
A1	0.26317	119	-12.8	DJ	Na	929	301	0.0 DJ, Lo
A1	525	031	-12.4	DJ	Na	929	303	- 2.0 Br, Gi,
A1	604	1.36998	-12.0	DJ		,,,	000	OL, Ruo,
Au	760	904	+19.2	ICT, FI				RZ, Wil
Cd	0.27486	637	+11.0	ICT, FI	Na	929	310	- 9.0 Ka
Hg	0.28035	442	+16.1	Ro	Na	929	308	- 7.0 Si
Αĺ	0.28163	428	-12.0	DJ	Na	929	293	+ 8.0 Du, Ket,
Hg	0.28936	168	+ 8.5	Ro			270	Rü, Wm
A1	0.30822	1.35671	+12.7	ICT, FI	Na	929	286	+ 15.0 Be
Al	822	694	-10.4	DJ	Na	929	280	4 21.0 La
Al	927	668	- 8.8	DJ	Hα	0.65628	115	
Hg	0.31317	567	+ 5.1	Ro				+ 0.3 ICT, OL, Ro, Sch
Hg Cd	0.33415	165	-1.6	Ro	Ħα	628	130	-14.7 Ka
Cd	0.34036	044	+16.0	ICT, FI	Hα	628	119	- 3.7 Br, Wil,
A1	0.35871	1.34795	-8.3	DJ				Wů
A1	0.36016	774	- 6.9	DJ	Hα	628	109	+ 6.3 Da, Du,
Çd	117	738	+15.8	IČT, FI		0.0	-07	La, Si
Al	124	760	- 7.1 - 2.6 + 7.3	DI	Hα	628	100	+15.3 Be
He	0.38886	432	- 2.6	Ro	He	0.66782	087	+ 0.4 Ro
Al	0.39440	366	+ 7.3	ICT, FI	Li	0.67079	079	+ 1.3 icr
A1	440	378	- 4.8	DJ	Li	079	082	- 1.7 Sch, Wm
Al	615	360	-3.8	DI	Li	079	087	- 6.7 Br
Ca	0.39685	1.34325	- 2.6	DJ ICT, Wil	Li	079	076	+ 4.3 Ket, Lo,
Ca	685	352	- 3.6	DJ			5,0	Rü
N	950	328	- 4.1	DJ	Li	079	073	+ 7.3 Du
Hg	0.40466	284	- 7.9	Ro	He	0.70652	003	- 3.1 Ro
N Hg Hδ	0.41017	228	- 1.0	Wil	K	0.76820	1.32888	-11.7 ICT
Ηδ	0.41017	208	+19.0	Du	K	820	884	- 7.7 Br, Sch
Fe	0.43258	029	+18.3	Du	ĸ	820	897	-20.7 Si
$H\gamma$	405	035	+ 1.6	ICT, Da		0.808	815	
$\Pi\gamma$	405	038	- 1.4	ICT, Da La, Sch		0.871	1.3270	- 11.1 Rub, Se - 2.6 ICT, Rub
$\Pi \gamma$	405	045	- 8.4	Br, Si		0.871	68	17.3 Se
Hγ	405	024	+12.6	Be		0.943	58	+ 3.5 ICT, Rub,
IIγ	405	015	+21.6	Du			30	Se Se
Hg	583	030	- 6.4	ICT, OL		1.000	1.323	+197. RL
Hg	583	027	- 3.4	Ro		1.028	1.3245	+ 4.9 ICT, Rub,
Hg	583	022 ₈	+ 0.8	Ja				Se Se
				-				

Table 137—(Continued)

λ	n	105(nc-	n) Ref.a	λ	n	$10^{3}(n-1.3)$	28) Ref. a
1.130	1.3230	+ 3.6	ICT, Ru	b. 12.0	1.187	-141	RL
1.150	1.0200		Se Se	13.0	1.269	-59	RL
1.256	1.3210	+14.8		ь. 15.0	1.332	+ 4	RL
1.200			Se	18.0	1.505	+177	RL
1.5	1.316	+214.	RL	25.5 to 26.	1.41d	+ 82	ICT, RH
1.617	1.3149		Se	46.9 to 53.6	1.36d	+ 32	ICT, RH
1.968	1.3078		Se	75.6 to 86.5	1.41 ⁴ 1.68 ⁴	+ 82	ICT, RH
2.0	1,300	+787.	RL	52	1.684	+352	ICT, RH CE CE CE CE
2.327	1.2997	+74.	Se	63	1.774	+442	CE
2.4	1.275	+2363.	RL	83	1.894	+562	CE
2.6	1,253	+4040.	RL	100	2.014	+682	CE
2.8	1.282	+574.	RL	117	2.044	+719	CE
3.0	1.365	-8335.	RL	152	2.094	+762	CE
				λ	n	(n-1.33)	Ref. a
λ	n	$10^{3}(n-1.328)$	Ref.				
3.0	1,365	+37	RL	4000.	9.50	8.17	Lam
3.2	1.456	+128	RL	4200.	5.33	4.00	ĮCT, T
3.4	1.437	+109	RL	6000.	9.40	8.07	Lam
3.6	1.384	+56	RL	8000.	8.97	7.64	Lam
3.8	53	+25	RI.	8400.	5.68	4.35	T T T
4.0	38	+10	RL	11000	6.27	4.94	T
4.5	43	+15	RL	15000	6.62	5.29	T
5.0	30	+ 2	RL	18000	6.65	5.32	T
5.5	1.300	-28	RL	27000	8.45	7.12	ICT, T
5.8	1.271	-57	RL				
6.0	1.324	- 4	RL	10 - 6x	n	(n-1.33)	Ref.ª
6.2	60	+32	RL	0.027	8.451	7.12	ICT, T
6.5	34	+ 6	RL	0.12 to 0.19	9.0	7.67	Šr ,
7.0	27	- 1	RL	0.375	9.08	7.75	Dr
8.0	1.293	-35	RL	0.5 to 0.6	Normale	7.73	2.
9.0	64	-64	RL	0.75	8.98	7.65	Dr
10.0	1.196	-32	RL	2.0	8.92	7.59	Dr
11.0	50	-178	RL	3.3 to 7.0	9.0	7.7	McCJ

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Table 137—(Continued)

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^b The $K\alpha$ line of characteristic x-radiation from Cu.

° At $\lambda = 0.1151 \,\mu$ the reflection is "metallic."

⁴ For $\lambda = 25.5$ to 152, the indicated values of *n* refer to the "residual" rays left after multiple reflection from the following solids: CaF₂, $\lambda = 25.5$ to 26; NaCl $\lambda = 46.9$ to 53.6; KBr, $\lambda = 75.6$ to 86.5; KCl, $\lambda = 63$; TlCl, $\lambda = 100$; TlBr, $\lambda = 117$, and TlI, $\lambda = 152\mu$. W. Weniger ⁴³² also, has found that in the interval $\lambda = 50 \mu$ to 300 μ the index is of the same order of magnitude as in the visible spectrum.

° It will be noticed that Lampa's values are much greater than those of T and ICT at $\lambda = 4200~\mu$ to $27000~\mu$. Lampa used the deviation by a prism, whereas Tear (T) derived n from the reflectivity and the extinction coefficient. Lampa concluded that the dispersion is normal in the range $\lambda = 8~\text{mm}~(=8000~\mu)$ to $1.2~\text{m}~(=12000000~\mu)$, but there is much absorption near $\lambda = 8000~\mu$, and hence, presumably, anomalous dispersion.

^t For $\lambda = 27000 \,\mu \, (=0.0270 \times 10^6 \,\mu)$, ICT gives n = 9.0.

"In the range $\lambda = 50$ to 60 cm (= 0.5 to 0.6 10" μ) there is no anomalous dispersion if the water is pure, that reported by R. Weichmann 4.51 being due to impurities, perhaps to dissolved glass.434

Table 138.—Refraction of Water at 20 °C: Comparison of Data

For the most accurate data, see Table 135; for other values, see Table 137 in which is given the formula by which the arbitrary norm (n_c) was computed. The preferred values are those in column TT.

Unit of $\lambda = 1 \mu = 10^4 \text{ A} = 10^{-4} \text{ cm}$. Index is with reference to air at 20 °C and 1 atm

Ref a→	TT	Be	Br	Da			Ket La						Wa	Wil	Wm	Wü
λ n_c							105(n	$n_{\epsilon})$								_
0.39440 1.34373 0.39685 1.34349						-7								+3		
0.43405 1.34037		-13	+7	-2	-22		+1				+1	+10		73		-1
0.44157 1.33983 0.48613 1.33716	-4	-11	+3	-11	-15	-2 +22	-4		-2		-1	-2		0		+1
0.53380 1.33498 0.53505 1.33493 0.58929 1.33301 0.65628 1.33115 0.67079 1.33080	2 0	-15 -15	-1 +3 +4 +7	-7	-11 - 9 - 6 - 7	+1 -1 +9 +15	-12 -10 -6	0 -2	+1 -1	-8 -7 -4	-2 -1 +1 +2		-2	+1 +2	-14 -9 +2	-6
0.76820 1.32876			+11								+6	+21				

a References:

TT Tilton, L. W., and Taylor, J. K., see Table 135; other symbols as in Table 137.

⁴⁸² Weniger, W., J. Opt. Soc. Amer., 7, 517-527 (1923).

⁴⁸⁸ Weichmann, R., Ann. d. Physik (4), 66, 501-545 (1921).

⁴³⁴ Mie, G., Physik. Z., 27, 792-795 (1926).

Other References.

For a discussion of procedures, instruments, and sources of error, see C. Chéneveau, 423 and L. W. Tilton 424; for a compilation of data prior to 1898, see H. Dufet 425; for a recent review of optical dispersion, see S. A. Korff and G. Breit 426; and for a discussion of the accurate representation of the index of refraction as a function of the wave-length and temperature, see L. W. Tilton 427 and L. W. Tilton and J. K. Taylor. 428

Table 139.—Dispersion Formulas for Water

$$(n_o)^2 = 1.762530 - 0.0133998 \lambda^2 + \frac{0.00630957}{\lambda^2 - 0.0158800} + \frac{10^{[107.731(0.064156 - \lambda^2) - 5]}}{10^{[107.731(0.064156 - \lambda^2) - 5]}}$$

$$(n_o')^2 = 1.762530 - 0.0133998 \lambda^2 + \frac{0.00630957}{\lambda^2 - 0.0158800}$$

$$(n_o'')^2 = 1.761480 - 0.0134140 \lambda^2 + \frac{0.00654380}{\lambda^2 - 0.0132526}$$

 n_c is used in Table 137; n_o' was proposed by Duclaux and Jeantet (see ftn., p. 280); and n_c'' by Martens and accepted by Chéneveau. n_o' is for 20 °C and n_o'' is for 18 °C. The index is with reference to air at the same temperature, at or near 20 °C; unit of $\lambda = 1 \mu$.

E. Flatow ⁴³⁵ represented his observations (2145A to 5893A) by means of 5 formulas of the type $n^2 = m - k \lambda^2 + m' \lambda^2/(\lambda^2 - \lambda_1^2)$, one for each temperature used. They may be put in the form $n^2 = a - b \lambda^2 + c/(\lambda^2 - \lambda_1^2)$, the values of the constants, referred to a vacuum, being these:

t	a	ь	c	$\lambda_{1^{2}}$
0	1.76565	0.013414	0.00633201	0.0159088
20	1.76362	0.013414	0.00626020	0.0161138
40	1.75758	0.013414	0.00619982	0.0162410
60	1.74840	0.013414	0.00613429	0.0163328
80	1.73755	0.013414	0.00603399	0.0165482

The unit of λ and of λ_1 is 1μ . The value for λ_1 at 20° as published in the original article and reproduced in compilations is $\lambda_1 = 0.12604 \mu$. That is obviously out of line with the others, and fails to reproduce the observations; it leads to c = 0.00617174 and $\lambda_1^2 = 0.158861$. It seems probable that the zero is a typographical error, and that the value should have been printed $\lambda_1 = 0.12694$. That leads to the values of c and λ_1^2 tabulated here, and fits the observations. Flatow's formulas are not considered further in this table. His values at 20 °C, referred to air, are in Table 137, where they are compared with n_c ; the mean temperature coefficients derived from them are in Table 141.

Very exact formulas based on their own observations have been given by L. W. Tilton and J. K. Taylor. 429

⁴⁸⁸ Flatow, E., Ann. d. Physik (4), 12, 85-106 (1903).

Table 139—(Continued)

 n_0 = observed value; values of $(n_0 - n_0)$ have been obtained, by interpolation, from Table 137; for $\lambda > 1.25$, $(n_0 - n_0)$ is great.

101 1

	Unit of $\lambda = 1 \mu$, $= 10^4 \text{ A}$; of $dn_0/d\lambda = 10^{-6} \text{ per A}$								
λ	n_o	$10^{8}(n'_{c}-n_{o})$	$10^6(n''_o - n_c)$	$10^{6}(n_{0}-n_{c})$	$-dn_c/d\lambda$				
(0.18) 0.19 0.20 0.22 0.25	(1.473230) 1.443823 1.423934 1.398714 1.377333	(-8979) -3652 -1406 -174 -5	(-23121) -12943 -7740 -3357 -1278	+3 +64 -15 +112	(362.6) 237.6 166.5 96.0 53.0				
0.30 0.40 0.50 0.60 0.70	1.358842 1.343192 1.336462 1.332683 1.330140	0 0 0 0	-333 -85 -132 -192 -236	-4 +41 +3 -6 +90	25.7 9.4 4.8 ₁ 3.0 ₀ 2.1 ₀				
0.80 1.00 1.25 1.50 2.00	1.328181 1.324969 1.321239 1.317272 1.307866	0 0 0 0	-271 -315 -348 -369 -397	+111 -42 -148	1.7 ₈ 1.5 ₀ 1.5 ₈ 1.6 ₇ 2.1 ₀				
2.50 3.00	1.296069 1.281653	0	-421 -446		2.6 ₂ 3.1 ₅				

Table 140.—Refraction of Water at Various Temperatures (See also Table 135.)

The values under HP and J have been derived from the corresponding formulas (see Table 142), the values beyond the range for which a formula is claimed to be valid being inclosed in parentheses. The others have been derived from the published experimental data. All the indices are with reference to air at the same temperature as the water.

Damien (Da) was positive that n continues to decrease as t decreases below zero; Jamin (J) stated that n is a maximum near zero, and with this the HP formula, resting on observations above 15 °C, agrees; see also text.

E. v. Aubel ⁴³⁶ has computed, on the basis of certain assumptions, that the index of refraction of water at the critical point is n = 1.102, probably for $\lambda = 6708$ A.

It will be noticed that the values of δ for the Ja observations exhibit surprising jumps at many of the places where the value of t was changed abruptly by several degrees.

 n_0 = value of n at t = 0; $n_0 - n = (n_0 - n)_{HP} + \delta$. For example, at -10 °C P's value for $10^5(n_0 - n)$ is 27, J's is 7; that is, P finds $n_{-10} = 1.33411 - 0.00027 = 1.33384$, and J finds $n_{-10} = n_0 - 0.00007$; J does not assign a value to n_0 .

⁴⁸⁶ v. Aubel, E., Physik. Z., 14, 302-303 (1913).

Table 140—(Continued)

Unit of $\lambda = 1A = 10^{-6}$ cm. Temp. = t° C

$ \begin{array}{c} \operatorname{Ref}^{a} \to \\ \lambda \to \\ n_{0} \to \\ t \end{array} $	HP 5893 D 1.33401 10 ^b (n _e -n) _{HP}	5893 D 1.33411	Da 6563 Ha 1.33225	Da 4862 Hs 1.33825 10%	Da 4341 Ηγ 1.34155	5893 D
-10 - 8 - 6 - 5	(22) (12) (6) (4)	+5 +4 +1 0	-20 -13	-19 -12	-19 -12	-15 -10 - 7 - 6
- 4	(2)	0	- 7	- 6	- 7	- 4
- 2 + 2	(0)	-1 +1	-3 + 2	- 2 + 3	- 3 + 2	- 2 + 3
T 4	(7)	T 1	+ 2	+ 3	+ 2	+ 3 + 1
5	(10)	+1				+ 1
6	(13)	+2	0	+ 1	+ 1	+ 1
8 10	(21) (31)	+1	$\begin{array}{ccc} + & 1 \\ + & 2 \end{array}$	$^{+11}_{+5}$	+ 2 + 4	+ 2 + 1
15	63	Ū	+ 2	+11	+ 4	- î
20	105		+12	+15	+15	- 3
$\operatorname{Ref}_{\substack{\lambda \to \\ n_0 \to }}^{a \to a}$	HP 5863 D 1.33401	1.334277	Hg 7 1	Ja 5770 Hg 1.334348	Ja 5461 Hg 1.335443	Ja 4358 Hg 1.341218
0.03	$10^{6}(n_{o}-n)$			10 ⁶ δ —	+ 1.8	
3.85	63.		23	- 23	- 19	- 16
5.71	120		20	20	- 29	- 22
5.76 6.55	122 151		32 37	- 30 - 35	- 24 - 32	- 20 - 23
6.63	154		•	00	- 29	- 21
7.88	206				- 32	- 23
8.09	216		37	- 35	- 33	- 29 - 23
8.52 8.85	235 252		31 42	- 44	- 39	- 23 - 27
9.15	266		37	- 36	- 34	- 24
9.44	281				- 36	- 28
9.65 14.06	292 563	_	66	- 63	- 34 - 53	- 25 - 34
15.00	631		65	- 62	- 56	- 34
15.24	649				- 62	- 42
15.96	703		72	- 70 105	- 60 - 93	- 50
21.44 22.19	1183 1256	- 1 -	09 97	- 105 - 97	- 93 - 85	- 61 - 48
23.20	1358	- 1		- 99	- 88	- 50
23.31	1370		99	- 97	- 85	- 48
24.42	1486 1534	- 1 - 1		- 103 - 99	- 88 - 84	- 51 - 45
24.87 27.67	1848	- i		- 109	- 97	- 51
28.16	1905	- 1	26	-114	- 104	- 71
28.60	1958	- 1		-114	- 99 103	- 51
28.65 29.25	1963 2035	-1 -1		-113 -116	- 103 - 105	- 65 - 49
39.51	3414		13		+ 8	+ 61
41.34	3688		21	- 7	+ 4	+ 64
47.45	4654		15	- 1	+ 13	+ 89
52.04	5433 7348		22 61	- 12 - 57	$^{+}_{-}$ 14 $^{-}$ 27	+ 107 + 99
62.42 75.95	10165		45	+ 38	+ 57	+287
89.63	13434					+288
92.25 93 .53	14109 14451		89 36	- 85 - 58	- 28	+278 +271

Table 140—(Continued)

* References:

Damien, B. C.⁴¹¹
Hall, E. E., and Payne, A. R., *Phys. Rev.* (2), 20, 249-258 (1922).
Jamin, J.⁴¹⁶
Jasse, C., *Compt. rend.*, 198, 163-164 (1934).
Pulfrich, C.⁴¹⁸ Da HP

Table 141.-Mean Temperature Coefficient of Index of Refraction of Water

(See also Table 135.)

In the lower right-hand corner of the table are given a number of values for the mean coefficient between 15 and 25 °C. Most of them have been derived from the formulas by which the several observers represent their observations.

The values in the rest of the table have been derived from observations, except those for C, HP, Ket, and R, which were computed.

 $\Delta t = (t_2 - t_1)$ °C; $\Delta n = n_2 - n_1$, n_1 and n_2 being the values of n at the temperatures t and t_2 °C respectively.

		Unit of $\lambda =$	$1A = 10^{-8} \text{ cm}$.	Γ emp. = t °C		
$t_1 \rightarrow t_2 \rightarrow \lambda$	20	20 40	40 60 10 ⁵ Δn/Δt	60 80	80 100	Ref.
2145	5.3	12.8	19.2	22.8_{5}		F1
2195	5.3 ₅	12.7	19.1	22.75		F1
2268	5.3	12.95	18.96	22.4		$\mathbf{F}\mathbf{l}$
2314	5.3_{5}	12.8	19.0	22.7		F1
2429	5.5	12.85	18.8	22.65		F1
2574	5.3	12.7	18.8	22.6		Fl
2677	5.3	12.5	18.6	22.5		F1
2749	5.25	12.6	18.55	22.35		F1
3082	5.0	12.45	18.4	22.18		F1
3404	4.9	12.3	18.3	21.95		Fl
3613	4.95	12.15	18.15	21.8		Fi
3945	4.75	12.2	18.05	21.65		F1
4341	6.0					Da
4417	4.65	12.1	17.9	21.55		Fl
4680	4.5₅	12.0	17.75	21.45		F1
4801	4.4	11.8	17.75	21.5		F1
4862	6.0					Da
5340	4.3	11.75	17.6	21.25		F1
5350	4.24	11.75	17.0	21.2	24.5	Ket
5350	4.13	11.85	17.75	20.8	19.75	R
5893	4.2	11.6	17.6_{5}	21.0		F1
5893	4.2	11.7	17.0	21.1	24.4	Ket
5893	5.25	12.2	16.95	21.0	25.75	HP
5893	4.1	12.0	17.4	21.05	•	C(ICT)
6563	5.85					Dà
6708	4.2	11.6	16.9	21.0	24.2	\mathbf{Ket}
6708	3.9	11.25	17.25	21.1	21.8	R

Table 141—(Continued)

Ref.a→ λ→	Ket 5350	0 T1R	Ket 670	R T.i	15 to 25 °C 5893	Ref.º
1		105(ns_			$-10^{6}\Delta n/\Delta t$	
10	19.1	20.8	18.9	19.6	9.17	HP
15	43.8	41.5	43.3	39.1	8.78	Wa
20	65.7	61.8	65.0	58.3	9.12	D
25	85.2	81.6	83.8	77.1	8.97	J
30	102	101	100	95	9.0	Wm
40	133	136	132	130		
50	160	166	158	160	7.93	F1
60	181	189	180	185	7.89	R
70	204	205	201	205	8.0	G
80	221	211	219	217		
90	236	206	234	221	8.4	Ket
100	255	189	251	215	8.4	Sch

a References:

Chéneveau, C., Int. Crit. Tables, 7, 12-16 (1930).

Dufet, H., Jour. de Phys. (2), 4, 389-419 (1885) ← Bull. soc. min. France, 8, 171-304 (1885).

Damien, B. C.⁴¹¹

Flatow, E.⁴⁸⁵

Gifford I. W. Proc. Roy. Soc. (London), (4), 78, 406-409, (1907). b

Da

- G HP
- Flatow, E. 435 (1907). Grant Payne, A. R., Phys. Rev. (2), 20, 249-258 (1922). Jamin, J. 416 Ketteler, E., Ann. d. Physik (Wied.), 33, 353-381, 506-534 (1888). Rühlmann, R., Idem (Pogg.), 132, 1-29, 177-203 (1867). Schutt, F., Z. physik. Chem., 5, 348-373 (1890). Walter, B., Ann. d. Physik (Wied.), 46, 423-425 (1892). Wiedemann, E., Idem (Pogg.), 158, 375-386 (1876). J Ket
- R Sch

Table 142.—Variation of the Refraction of Water with the Temperature: Comparison of Formulas

In the first section of the table are collected formulas that are quoted in one or more widely used compilations or that have been proposed recently; in the second is a skeleton table comparing the values defined by those formulas for the D-lines (5893A). Key letters indicate the several formulas and their sources. Values beyond the range in t assigned to a formula are enclosed in parentheses.

Flatow did not give a formula c connecting n and t, but gave five dispersion formulas, one for each of the temperatures used (see Table 139.)

Unit of v^* , B, and C=1 cm³ g; of k=1 per 1 °C; of $\lambda=1$ A=10-8 cm. Temp. = t °C

Formulas for the variation of n with t.

Ketteler's formula $(n^2 - 1)(v^* - \beta) = C(1 + \alpha e^{-kt})$, where α , β , and k are constants determined solely by the material, v^* is the specific volume, and C depends upon the substance and the frequency of the radiation, is in disagreement with Flatow's observations (see text), and the present compiler has been unable to check Ketteler's computations satisfactorily. Possibly the values of v* used by Ketteler in deriving the constants, given below, were unsatisfactory at the higher temperatures.

Very precise formulas based on their own observations have been published by L. W. Tilton and J. K. Taylor. 437

⁴⁸⁷ Tilton, L. W., and Taylor, J. K., J. Res. Nat. Bur. Stand., 20, 419-477 (RP1085) (1938).

Table 142—(Continued)

Other	formulas	are of	the typ	$e 10^5(n-n_0)$	$= at + bt^2$	$+ ct^3 + dt^4$
-------	----------	--------	---------	-----------------	---------------	-----------------

Key [∉]	102a	10°b	105c	10°d	Range in t	λ	Basis ^b
C°	-12.4	-199.3	0	+5.	0 to 80	(D)5893	Air, t
\mathbf{D}	-125.5	-206.4	+4.35	+11.5	1 to 50	(D)5893	Air, R
HP^d	- 66.	-262.	+181.7	- 7.55	15 to 100		Air, 20
J	-125.73	- 192.9	0	0	0 to 30	(D)5893	Vac
Lo	+0.76	-280.3	+213.4	0		(D)5893	Vac
P	- 83.	- 295.	+640.	0	-10 to 10	(D)5893	Air, t
\mathbf{M}^{f}	-20.0	-290.5	0	+5.00		(D)5893	•
R	0	-201.4	0	+4.936		(D)5893	Air, 9
Wa	- 120 .	- 205 .	+50.	0	0 to 30	(D)5893	Air, $R(?)$
Lo	+9.52	-279.3	+213.4	0	0 to 30	(Li)6708	Air, $R(?)$
R	0	-196.6	0	+4.600	0 to 92	(Li)6708	Air, 9
R	0	-209.0	0	+6.046	0 to 92	(T1)5350	Air, 9
О	- 118.73	- 207.09	+7.92	+10.939	2 to 38	(Hg)5461	Vac
Keya	105α	105β	10 ⁵ C	105k	Range in t	λ	Basis ^b
Ket	246	20271	61574	2290	0 to 100	(Li)6708	Vac
Ket	246	20271	62035	2290	0 to 100	(D)5893	Vac
Ket	246	20271	62439	2290	0 to 100	(T1)5350	Vac

II. (D) $\lambda = 5893A$: Comparison of formulas.

For simplicity, all formulas have been compared with the most recent one (HP); $10^5(n_0 - n_t)$ is given for the HP formula, and that plus δ is the corresponding quantity for the other indicated formula; $(n_0 - n_t)_{\rm vac} = (n_0 - n_t)_{\rm air} + 10^{-5}\Delta$; the tabulated values of Δ have been derived from Table 136.

Ref ^e Date Basis ^b	HP 1922 Air, 20	Wa 1892 Air, <i>R</i>	D 1885 Air, <i>R</i>	J 1856 Vac	P 1888 Air, <i>t</i>	Lo 1880 Vac	C 1930 Air, t	R 1867 Air, 9	Ket 1880 Vac	Δ
t	$10^5(n_0-n_i)$)				8				
-10	(22)				6					-1.5
- 5	(4)				0					-0.8
+5	(10)	1	1	1	1	-3	-4	- 5	-7	+0.7
10	(31)	1	2	1	0	-5	-10	- 10	- 12	1.4
15	63	- 1	1	- 1	(-6)	-7	-17	-18	-17	2.0
20	105	-3	1	- 2		-10	-23	-25	-20	2.7
25	155	-4	0	-3		-13	- 29	-31	-25	3.3
30	213	-6	0	- 1		-18	-34	- 35	-28	3.9
40	349	(-5)	$-\frac{0}{7}$	(+10)		(-37)	-37	-38	-31	5.0
50	509		-7				- 35	- 36	-32	6.1
60	688		(-28)				-28	- 27	-30	7.2
70	887		,				-22	- 19	-27	8.1
80	1109						-28	- 22	-28	9.1
90	1352						(-55)	-45	-37	9.9
100	1624							(-104)	-56	10.8

* References:

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C Same as Table 141.
D Same as Table 141.
HP Same as Table 141.
J Jamin, J. 616
Ket Ketteler, E., Ann. d. Physik (Wicd.), 33, 353-381, 506-534 (1888).
Lo Lorenz, L. 617
O Osborn, F. A., Phys. Rev. (2), 1, 198-210 (1913).
P Pulfrich, C. 618
Rühlmann, R., Ann. d. Physik (Pogg.), 132, 1-29, 177-203 (1867).
Wa Walter, B., Idem (Wied.), 46, 423-425 (1892).
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Table 142—(Continued)

Basis: (Air, t), (Air, 9), (Air, R), etc., and (Vac) indicate that the associated n is that with reference to air at the temperature (t) of the water, at 9 °C, at room temperature, etc., and with reference to vacuum, respectively. Quite frequently observers and compilers state that a value of n refers to "air at the same temperature (Air, t) when it actually refers to air at room temperature (Air, R) (see citations R and HP). Here the basis is given as (Air, R) unless there is definite evidence that it is some other. In all cases, the pressure of the air of reference is 1 atm.

The difference $(n_0 - n_1)$ when referred to vacuum (Vac) exceeds its value when referred to air at a fixed temperature (Air, R; Air, 9; Air, 20) by only 3 parts in 10 000, an amount that is negligible when, as here, the index is not carried beyond the fifth place of decimals. But when the index is with reference to air at the same (varied) temperature as the water (Air, t), then the value of $(n_0 - n_t)$ for (Vac) does, in general, differ significantly from that for (Air, t).

*The original source of this formula (C), given in Chéneveau's compilation, has not been ascertained. The Landolt-Börnstein Tabellen attributes it to F. F. Martens, who prepared that section of the Tabellen, and states that it is based on Flatow's observations, but it is not given in any of Martens' papers that have come to the attention of the compiler.

^d In their synopsis, the authors state that the values of n when referred to vacuum are represented by the formula: $n = 1.33401 - 10^{-7}(66t + 26.2t^2 - 0.1817t^3 + 0.000755t^4)$, but in reality that formula, which agrees with the one here given, reproduces their values as referred to air, and given in their Table II; it does not reproduce their vacuum values, given in Table IV.

This formula seems to have been derived by H. Dufet 485 from the observations of Pulfrich.

The Landolt-Börnstein Tabellen attributes this formula to F. F. Martens, who prepared that section of the Tabellen, and states that it is based on the data of Pulfrich, but it does not satisfactorily represent those data and has not been found in Martens' papers. It does not appear elsewhere in this compilation, and is not used in the following sections of this table.

Table 143.—Temperature Gradient of the Index of Refraction of Water

(See also Table 135.)

In the first section, the values of dn/dt for $\lambda = 5893$ A are given as derived from the several formulas listed in Table 142; the key symbols are the same in both tables. In the second, experimental values for 20 °C and various λ 's are given.

If n is with reference to air at a fixed temperature (Air, R; Air, 9; Air, 20), then dn/dt has essentially the same value (within 3 parts in $10\,000$) as if n had been referred to a vacuum; such is not the case if n is with reference to air at the same (varied) temperature as the water (Air, t).

For $\lambda = 12.6$ to 24 cm, M. Seeberger ^{437a} has concluded that $-10^5 dn/dt =$ 1600, essentially independent of λ and of the temperature throughout the range 16 to 70 °C.

⁴⁸⁷a Seeberger, M., Ann. d. Physik (5), 16, 77-99 (1933).

Table 143—(Continued)

I. Temperature gradient as derived from formulas. $\lambda = 5893 A(D)$. $-10^{5} dn/dt = -10^{5} (dn/dt)_{HP} + \delta$. Temp. = $t \, ^{\circ}$ C.

$Ref.^{a} \rightarrow Basis^{b} \rightarrow t$	HP Air, 20 105dn/dt	Wa Air, R	D Air, R	J Vac	P Air, t —— 100δ —	Lo Vac	C Air, t	R Air, 9
- 10	(-5.12)	(+207)	(+99)	(+252)	- 186	(-113)	(+134)	(+111)
-5	(-2.10)	(+121)	(+97)	(+143)	- 50	(-87)	(+23)	(+9)
0	(+0.66)	+54	+60	+60	+17	- 67	- 54	- 66
0 +5	(3.15)	+6	+16	+4	+15	- 52	- 103	- 114
10	(5.38)	-23	-6	- 26	-57	-42	- 129	- 137
15	7.40	-39	- 14	-36	(-204)	- 44	- 136	-142
20	9.20	-40	-11	-23	,	- 56	- 126	- 130
25	10.82	-31	-4	+8		-81	-104	- 106
30	12.29	- 14	- 1	+54		-124	-75	-74
40	14.83	(+37)	-21	(+186)		(-266)	-4	+2
50	17.01		-119				+54	+66
60	19.00		(-338)				+72	+90
70	20.99		,				+18	+43
80	23.16						- 139	-104
90	25.68						(-426)	-382
100	28.75						-	-821

II. Experimental values at 20 °C.

λ	1862	1930	1988	1990	2144	2144	5893A
$-10^5 dn/dt$	8.2	9.3	6.9	10.5	8.90	7.2	8.0
Basis ^b	Air, 20	Air, 20	Air, 15	Air, 20	Air, 20	Air, 15	Air, 15
Ref.d	DI	DJ	Ġ	DI	DJ	G	Ġ

[&]quot;As in Table 142.

Table 144.—Variation of the Refraction of Water with the Pressure

$$\Delta \equiv (n_2 - n_1)/(p_2 - p_1); dn/dp = a - bp.$$

G. Quincke 438 did not accept Zehnder's values (see below), believing that his own observations 439 indicated that when the temperature is constant, then $(n-1)/\rho$ is a constant, ρ being the density.

For effect of pressure on dielectric constant see Table 176 and accompanying text.

Unit of Δ and of $a=10^{-6}$ per atm; of $b=10^{-6}$ per atm²; of $\lambda=1$ A = 10^{-8} cm. Temp. = t °C

I. D-lines, $\lambda = 5893$. Pressure not exceeding 4 atm.

Mascart ⁴⁴⁰ found $\Delta = 15.2$ at 15 °C, and 16.1 at 5.5 °C.

The following observations (Δ_{obs}) from L. Zehnder ⁴⁴¹ are closely reproduced by $\Delta_c = 16.84 - 0.129t + 0.0022t^2$.

^b Basis of reference, see Table 142. ^c This value is attributed by DJ to E. Flatow. ⁴³⁵ DJ conclude that -dn/dt passes through a maximum near $\lambda = 2000 A$.

^d References:

DJ Duclaux, J., and Jeantet, P.⁴¹⁵ G Gifford, J. W., Proc. Roy. Soc. (London) (A), 78, 406-409 (1907).

⁴³⁸ Quincke, G., Ann. d. Physik (Wied.), 44, 774-777 (1891).

⁴⁸⁹ Quincke, G., Idem, 19, 401-435 (1883).

⁴⁴⁰ Mascart, Compt. rend., 78, 801-805 (1874).

⁴⁴¹ Zehnder, L., Ann. d. Physik (Wied.), 34, 91-121 (1888).

Table	144	(Continued)
Table	144	(Continuea)

t	Δ_{obs}	Δ_c	t	Δ_{obs}	Δe
-0.78	16.91	16.94	4.95	16.26	16.26
0.00	16.82	16.84	8.95	15.87	15.86
+0.06	16.85	16.83	9.00	15.91	15.86
0.42	16.78	16.79	13.05	15.55	15.53
1.05	16.69	16.71	13.28	15.56	15.52
2.62	16.51	16.52	17.83	15.25	15.24
2.67	16.53	16.51	18.01	15.25	15.23
2.92	16.48	16.48	18.03	15.25	15.23
3.10	16.44	16.46	23.27	14.98	15.03

II. Various wave-lengths.

Low pressur	es; 18 °C RZª	Pressure	up to 1800 kg/cm ²	: 25 °C PRª
λ	Δ	λ	a	b
4861	15.40	4060	15.02	0.003182
6807	15.16	4360	14.65	0.002700
		5460	14.75	0.003132
		5790	14.56	0.002990

a References:

PR Poindexter, F. E., and Rosen, J. S., Phys. Rev. (2), 45, 760 (A) (1934). RZ Röntgen, W. C., and Zehnder, L., Ann. d. Physik (Wied.), 44, 24-51 (1891).

Table 145.—Refraction of Natural Waters

 $n = n_w + \Delta$, n_w being the index for pure water at the same temperature and for the same wave-length; salt content = s g per kg of sea-water.

C. Chéneveau a gives the following values for 20 °C and the D-lines:

	10-4
City water, Paris	4
River Seine	5
Water saturated with CO ₂ at 1 atm	-3
Mediterranean Sea	400

The last seems to be entirely too small; see below.

Unit of $dn/dt = 10^{-6}$ per 1 °C; of s = 1 g salt per kg sea-water; of $\lambda = 1$ A = 10^{-8} cm

		Sea-water -			
—J. W. Gifford (1907)		Mediterranean Sea 20° C		_O. Krummel•_ D-line; 18 °C	
15° C					
10⁵∆	-dn/dt	λ	10 ^δ Δ	s	$10^5(n_o-n_w)$
647		(A) 7608	697	5	97
645		(B) 6870	691	10	194
650		(C) 6563	696	15	290
653	78.5	(D) 5893		20	386
664		(F) 4862			482
665					577
676		(H) 3969	756		673
680				40	769
691					
698		$n_{10} - n_{20} = 0$.00085		
819					
	75.8				
Abs					
	15° C 10° Δ 647 645 650 653 664 665 676 680 691 698 819 979	15° C 10° \(\triangle \triangle -dn/dt \) 647 645 650 653 78.5 664 665 676 680 691 698 819 74.7 979 75.8	fford $(1907)^a$ SS (1889) 15° C 20° C $10^0\Delta$ $-dn/dt$ λ 647 (A) 7608 645 (B) 6870 650 (C) 6563 653 78.5 (D) 5893 664 (F) 4862 665 (h) 4102 676 (H) 3969 680 691 698 $n_{10} - n_{20} = 0$ 819 74.7 979 75.8	fford $(1907)^a$ — SS $(1889)^a$ — Mediterranean Sea 20° C 20° C $10^{\circ}\Delta$ — dn/dt	fford $(1907)^a$ — SS $(1889)^a$ — O. Kr Mediterranean Sea 20° C D-lin $10^{\circ}\Delta$ — dn/dt λ 10° Δ s 647 (A) 7608 697 5 645 (B) 6870 691 10 650 (C) 6563 696 15 653 78.5 (D) 5893 706 20 664 (F) 4862 719 25 665 (h) 4102 739 30 676 (H) 3969 756 35 680 691 691 698 819 74.7 979 75.8

^b This value accords with the published coefficient, but seems strangely out of line with the others.

Table 145—(Continued)

" References:

Chéneveau, C., Int. Crit. Tables, 7, 12-16 (13) (1930) from observations by Dufet, H., Bull. Soc. Min. France, 8, 171-304 (1885); Soret, J. L., and Sarasin, E., Compt. rend., 108, 1248-1249 (1889).
 Gifford, J. W., Proc. Roy. Soc. (London) (A), 78, 406-409 (1907).
 Krümmel, O. 608

SS Soret, J. L., and Sarasin, E., Compt. rend., 108, 1248-1249 (1889) quoted by Krümmel. Strong absorption for $\lambda \ge 2194.4$ A.

REFLECTION OF LIGHT BY WATER

When light strikes a boundary separating two media of different refractivities, some of it is specularly reflected, some is scattered (nonspecularly reflected), and the remainder enters the second medium. amount that is specularly reflected depends upon both the angle of incidence (i) and the polarization of the incident light. Unless i = 0, the reflected light is partially plane-polarized even when the incident light is not, for the component that has its electric vector perpendicular to the plane of incidence (i.e., that is polarized in the plane of incidence) is more strongly reflected than the other. As i increases from 0 to 90°, the ratio of the reflectivities of the two components passes through a maximum. angle at which this occurs is known as the Brewsterian angle and is given by the relation $\tan i = n$. At that angle the reflected light is almost completely plane-polarized, the reflectivity of the weaker component, in the case of an air-water surface, being only a few ten-thousandths of that of the stronger; C. V. Raman and L. A. Ramdas 442 found for that ratio 75×10^{-5} , while Rayleigh ⁴⁴³ found 42×10^{-5} . From their own measurements, Raman and Ramdas computed that the thickness of the transition layer in which the index changes from that of air in bulk to that of water in bulk is of the order of 5×10^{-8} cm; the diameter of a water molecule, as calculated from viscosity data, is about 2.6×10^{-8} cm.

The reflectivity (R) is defined by the ratio $R = I_r/I_4$ where I_4 and I_r are the intensities of the incident and of the specularly reflected light, respectively. If the medium is transparent (both absorption and scattering negligible), the reflectivity is $R_p = \frac{\sin^2(i-r)}{\sin^2(i+r)}$, if the incident light is plane-polarized in the plane of incidence (electric vector parallel to the reflecting surface); and $R_n = \frac{\tan^2(i-r)}{\tan^2(i+r)}$ if it is plane-polarized normal to the plane of incidence; i is the angle of incidence and r that of refraction. If the incident light is unpolarized, the reflectivity is half the sum of these

two expressions. As the index of refraction of the second medium with

⁴⁴⁹ Raman, C. V., and Ramdas, L. A., Phil. Mag. (7), 3, 220-223 (1927).

⁴⁴⁸ Lord Rayleigh, Phil. Mag. (5), 33, 1-19 (1892)="Collected Works," Vol. 3, pp. 496-512, 1902.

reference to the first is $n = \sin i/\sin r$, these expressions may be put in the form:

$$R_p = \frac{[\sqrt{n^2 - \sin^2 i - \cos i}]^2}{[\sqrt{n^2 - \sin^2 i} + \cos i]^2} \text{ and } R_n = \frac{[n^2 \cos i - \sqrt{n^2 - \sin^2 i}]^2}{[n^2 \cos i + \sqrt{n^2 - \sin^2 i}]^2}$$

If i = 0, $R_p = R_n = (n-1)^2/(n+1)^2$. Throughout the visible spectrum the transparency of water is such that R for it can be satisfactorily computed by means of these formulas. For experimentally determined values of R see Table 146.

The scattering (non-specular reflection) of light by a free liquid surface probably arises from the roughening of the surface by the thermal agitation of the molecules. The higher the surface tension and the more nearly

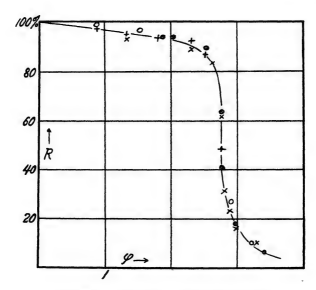


FIGURE 4. Reflection of X-rays by Water.

[Adapted from H. Steps, Ann. d. Physik (5), 16, 949-972 (1933).] Radiation used was the $K\alpha$ of Cu (λ = 1.539A); R = reflectivity; 90° – ϕ angle of incidence, unit - 0.001 radian = 0.057° = 3.44′.

equal are the indices of refraction of the adjacent fluids, the less the scattering. The observed ratio of the intensity of the light scattered by an airwater surface at room temperature to that scattered by plaster of Paris varies from 3×10^{-6} to 8×10^{-6} .

V. E. Shelford and F. W. Gail 445 have reported that in calm, clear weather between 10 a.m. and 2 p.m. about 25 per cent of the light from the

⁴⁴⁴ Raman, C. V., and Ramdas, L. A., Proc. Roy. Soc. (London) (A), 109, 150-157, 272-279 (1925). See also Ramdas, L. A., Indian J. Phys., 1, 199-234 (1927).
445 Shelford, V. E., and Gail, F. W., Publ. Puget Sound Biol. Sta., 3, 141-176 (1922).

Table 146.—Reflectivity of Water

Observed values for unpolarized light. For the reflectivity of polarized light, see remarks in text; of x-rays, see Fig. 4.

Except as noted, the data have been taken from E. P. T. Tyndall's compilation ⁴⁵⁰ based on data obtained by A. K. Ångström, ⁴⁵¹ K. Brieger, ⁴⁵² F. Gehrts, ⁴⁵³ H. Rubens, ⁴⁵⁴ H. Rubens and E. Ladenburg. ⁴⁵⁵

More recent data obtained by M. Weingeroff ⁴⁵⁶ and published as a small scale curve covering the range $\lambda = 11 \,\mu$ to $\lambda = 17 \,\mu$ essentially agree with these except at the longer wave-lengths, where he finds a somewhat smaller reflectivity.

 $R = I_r/I_i$, where I_i and I_r are the intensities of the incident and the reflected light, respectively; i is angle of incidence; $\lambda_{\max} = \text{wave-length}$ at which R passes through a maximum.

		Unit of $\lambda = 1 \mu = 10$	-4 cm. Temp. = 2	0 °C	
3.2 3.23		4.7	6.3 6.22	19.5	Ref.ª RL Re
3.045, 3.15, 3.28(?)			6.40		BM
3.18	3		6.40		MB1
	100R		I Incidence	λ	100R
λ		λ	100R		
0.3	2.33	6.3	2.34	18.0	6.7
0.7	2.00	6.5	2.10	18.5	7.5
1.0	1.98	7.0	1.95	19.0	8.4
1.5	1.95	7.5	1.75	20.0	8.9
2.0	1.74	8.0	1.67	21.0	8.2
2.4	1.45	8.5	1.60	23.	6.50
2.6	1.25	9.0	1.44	33.	7.2°
2.8	1.35	9.5	1.24	52.	9.30
3.0	3.40	10.0	0.95	52.	9.30^{d}
3.2	4.10	10.5	0.85	63.	10.60
3.23	3.4	11.0	0.75	63.	10.74^{d}
3.4	3.25	11.5	1.10	82.	9.6
3.5	2.95	12.0	2.00	83.	10.90
4.0	2.20	12.5	2.00	83.	11.75d
4.5	2.14	13.0	3.10	94.	11.10
5.0	2.00	13.5	3.40	100.	12.28d
		14.0	4.10		
5.5	1.68			108.	11.6
5.6	1.67	14.5	4.80	117.	12.80^{d}
5.8	1.40	15.0	5.30	117.	12.70
5.9	1.50	15.5	5.4	152.	13.40^{d}
6.0	2.00	16.0	5.3	310.	15.10
6.1	2.28	16.5	6.0	4200.	48.4
6.2	2.46	17.0	6.6		
6.22	2.4	17.5	6.9		
Normal	Incidence		Un	$\text{nit of } \lambda = 1\mu$	
Timit of	λ=1 cm-			i = 50°	
λ Unit of	100 R	λ	100 R	λ	100 R
		6.0	4.00	13.0	4.70
0.42	48.4	7.0	3.25	14.0	6.65
0.84	51.6 ^f	8.0	3.20	15.0	8.20
1.1	54.1	9.0	2.50	16.0	8.7
1.5	56.51	10.0	2.10	17.0	9.8
1.8	58.1	11.0	1.80	18.0	11.3
2.7	64.1	12.0	2.80	19.0	13.7

Table 146—(Continued)

Effect of temperature; normal incidence Unit of $\lambda = 1 \mu = 10^{-4}$ cm. Temp. = t° C

$t \rightarrow \lambda$	0	20 100R ——	30
117	11.8	12.7	13.2
310	14.9	15.1	17.1

* References:

Barnes, R. B., and Matossi, F., Z. Physik, 76, 24-37 (1932).

Matossi, F., and Bluschke, Idem, 104, 580-583 (1937). See also Matossi, F., and Fesser, H., Idem, 96, 12-28 (1935).

Reinkober, O., Idem, 35, 179-192 (1926).

Rubens, H., and Ladenburg, E., Verh. physik. Ges., 10, 226-227 (1908). MB1

^b Calculated from n.

° R. Rubens *; previously, he and H. Hollnagel † had found for the residual rays from KBr ($\lambda = 74$ to $88\,\mu$) 100R = 9.6 at 19 °C, R for silver being taken as unity.
⁴ C. H. Cartwright and J. Errara.‡
⁵ H. Rubens and R. W. Wood.§

'J. D. Tear,** radiation was plane-polarized with the electric vector in the plane of incidence; $i = 8^{\circ} 20'$.

Table 147.—Albedo of Water 157

By definition, the albedo of a plane surface is $A = F_r/F_t$ where F_r is the total luminous flux reflected by the surface when uniformly illuminated by white light, the total luminous flux incident on the surface being F_i .

The following values were taken from an airplane at altitude H, and in some cases the reflected light passed through one or other of two color filters described simply as "green" and "red," respectively.

Unit of H = 1 ft = 0.3048m; of $A = 10^{-8}$

$\underset{H}{\text{Filter}} \rightarrow$	None	Green A	Red	Remarks
	·	Chesapeake Bay		
2000	97	onionpinnio Luj		Smooth
2000	384	47	35	Well out
3000	36	40	45	Well out
		Potomac River		
10 to 20	69	55	104	
		Patuxent River		
2000	55			
3000	64			

^a At Buzzards Bay, W. M. Powell and G. L. Clarke ⁴⁸⁸ found A=3 to 4 per cent.

^{*} Rubens, R., Sitzb. preuss. Akad. Wiss. (Berlin) (Phys.-Math.), 1915, 4-20 (1915).
† Rubens, R., and Hollnagel, H., Ber. physik. Ges., 12, 83-98 (1910).
‡ Caitwright, C. H., and Errera, J., Proc. Roy. Soc. (London), (A). 154, 138-157 (1936); Acta Physicochim. URSS, 3, 649-684 (1935)→Cartwright, C. H., Nature, 135, 872 (L) (1935).
§ Rubens, H., and Wood, R. W., Verh. physik. Ges., 13, 88-100 (1911).
** Teat, J. D., Phys. Rev. (2), 21, 611-622 (1923).

⁴⁴⁰ Hulburt, E. O., J. Opt. Soc. Amer., 24, 35-42 (1934).

⁴⁴⁷ Weniger, W., J. Opt. Soc. Amer., 7, 517-527 (1923).

⁴⁴⁸ Schaefer, C., and Matossi, F., "Das Ultrarote Spektrum," Berlin, J. Springer, 1930.

⁴⁴⁹ Korff, S. A., and Breit, G., Rev. Mod Phys. 4, 471-503 (1932).

⁴⁵⁰ Tyndall, E. P. T., Int. Crit. Tables, 5, 258-259 (1929).

⁴⁵¹ Angström, A. K., Phys. Rev. (2), 3, 47-55 (1914).

⁴⁵² Brieger, K., Ann. d. Physik (4), 57, 287-320 (1918).

⁴⁵⁸ Gehrts, F., Idem, 47, 1059-1088 (1915).

sky is reflected by the surface of the sea; and E. O. Hulburt 446 has found that, when the sky is clear, the brightness of the rim of the sea is 25 per cent of that of the sky near the horizon, the surface of the sea being ruffled by a breeze of 5 to 25 knots, under which condition, "The reflecting facets of the sea which are visible to the observer are tilted up on an average of 15° from the horizontal." He defines the rim of the sea as the surface lying between the horizon and a line of sight making an angle of 3° with the surface of the sea.

For reviews and summaries see W. Weniger,⁴⁴⁷ C. Schaefer and F. Matossi,⁴⁴⁸ S. R. Korff and G. Breit.⁴⁴⁹

39. LUMINESCENCE OF WATER

Luminescence may be excited in water by various means: by light, x-rays, and gamma rays, by bombardment with electrons and beta rays, and by mechanical shock. These will be considered in the order named, after certain terms have been defined and certain general characteristics of the several types of luminescence excited by light have been briefly considered.

Types of Luminescence. Definitions and General Characteristics.

When a beam of light is passed through a medium, the medium becomes luminous, emitting light even at right angles to the incident beam. This luminescence may be very faint, and is observed most satisfactorily in a direction at right angles to the incident beam. It is usually partially polarized, even when the exciting light is unpolarized. The intensity of the component polarized in a given plane varies with the orientation of that plane about the line of sight, passing through a maximum (I_s) and a minimum (I_w) . The ratio $\rho = I_w/I_s$ is called the depolarization factor, and $\Delta = 2I_w/(I_w + I_s) \equiv 2\rho/(1 + \rho)$ is called the depolarization. These two quantities should not be confused.

Four distinct types of such luminescence are recognized: Fluorescence and phosphorescence, Tyndall scattering, Rayleigh scattering, and Raman scattering.

Fluorescence and Phosphorescence. — Phosphorescence being merely long-lived fluorescence, the latter, and shorter, term will be used for both. Fluorescence differs from other types of luminescence in that the spectrum of its light depends solely upon the medium, and in that the light is not polarized except as polarization may be imposed upon it by the incident radiation. But any specified portion of the spectrum of the fluorescent light may appear only under certain conditions; e.g., only when the exciter contains frequencies lying within a certain range, or exceeding a certain

⁴⁶⁴ Rubens, H., Verh. physik. Ges., 17, 315-335 (1915).

⁴⁶⁵ Rubens, H., and Ladenburg, E., Idem, 11, 16-27 (1909); Sitz. preus. Akad. Wiss, 1908, 274-284 (1908).

⁴⁵⁶ Weingeroff, M., Z. Physik, 70, 104-108 (1931).

⁴⁸⁷ Kimball, H. H., and Hand, I. F., Monthly Weather Rev., 58, 280-281 (1930).

⁴⁵⁸ Powell, W. M., and Clarke, G. L., J. Opt. Soc. Amer., 26, 111-120 (1936).

value, or contains corpuscles having a kinetic energy exceeding a certain value

Tyndall Scattering.—If small foreign particles are distributed throughout the medium, they will scatter the light by reflection and diffraction. The spectrum of the light so scattered (Tyndall scattering) is the same as that of the incident light as modified by the color of the scattering particles. If they are colorless, the ratio of the intensity of the scattered to that of the incident light varies continuously throughout the spectrum, the rate of variation at any place depending upon the size of the particles, as well as upon the wave-length. If the particles are very small, the ratio varies inversely as the fourth power of the wave-length of the light, making the scattered light much bluer than the incident. If the incident light is unpolarized, the light so scattered at right angles to the incident beam will be completely plane-polarized in the plane of scattering (the electric vector being normal to that plane) if the particles are spherical and isotropic; otherwise, the polarization will not be complete.⁴⁵⁹

Ravleigh Scattering.—An exactly analogous scattering by pure, dustfree gases was predicted by Lord Rayleigh 400 and has been observed. It arises from the scattering by the molecules themselves, which here play exactly the same part as is played by the foreign particles in the Tyndall scattering. It is but a step to extend the same idea to liquids and solids, but in them the molecules are so closely packed that they cannot satisfactorily play the part of foreign particles. Nevertheless, liquids and solids do exhibit exactly this same type of scattering, the scattering "particles" in them being the slight variations from point to point, and from instant to instant, in the number of molecules per unit of volume, these variations arising from the thermal agitation of the molecules.⁴⁶¹ Somewhat similar variations in the concentration of the primary molecules (H2O) may arise from the temporary association of these molecules into rather large groups, as suggested by H. Schade and H. Lohfert 162 and by G. W. Stewart, 463 who uses the adjective cybotactic to describe such a condition of association. It would seem that such groups also might act as scattering particles.

Scattering of this type by pure, dust-free media, whether liquid, solid, or gas, will be called Rayleigh scattering. To the light so scattered applies everything that has been said about Tyndall scattered light, except that the scattered light will never be completely polarized if there is interaction between the scattering units, no matter how nearly isotropic the units may be. Each type of scattering—Tyndall and Rayleigh—is frequently called by either name.

⁴⁵⁰ See Strutt, J. W. (later, Lord Rayleigh), Phil. Mag. (4), 41, 107-120, 274-279 (1871); 41, 447-457 (1871). Lord Rayleigh, Idem (5), 12, 18-101 (1881); (6), 35, 373-381 (1918).

400 Lord Rayleigh, Phil. Mag. (5), 47, 375-384 (1899).

⁴⁰¹ See v. Smoluchowski, M., Ann. d. Physik (4), 25, 205-226 (1908); Einstein, A., Ann. d. Physik (4), 33, 1275-1298 (1910); Raman, C. V., Proc. Roy. Soc. (London) (A), 101, 64-80 (1922).

⁴⁰² Schade, H., and Lohfert, H., Kolloid Z., 51, 65-71 (1930).

⁴⁰⁰ Stewart, G. W., Phys. Rev. (2), 35, 726-732 → 1426 (A) (1930).

⁴⁶⁴ Cabannes, J., Jour. de phys. (6), 3, 429-442 (1922).

⁴⁶⁸ Cabannes, J., and Daure, P., Compt. rend., 186, 1533-1534 (1928).

In 1928 J. Cabannes and P. Daure 465 announced that the radiation in the Rayleigh scattered light appeared to have a slightly lower frequency (displacement of about 0.01A toward the red), and that the line appeared broader than in the incident light and was superposed on a rather sharply limited continuous background, the whole having the appearance of a winged line. This last is sometimes referred to as the Cabannes-Daure effect. The work was continued by J. Cabannes and P. Salvaire, 406 J. Cabannes, 467 and others; and more recently by W. Ranin, 468 who found that when the incident radiation is truly monochromatic the scattered line is a symmetrical triplet, of which the central line has, within experimental error, the same frequency as the incident radiation. All three components of the triplet are of about the same intensity, and the spacing agrees well with that called for by L. Brillouin's theory of scattering. 469 The triplet rests on a continuous background, as Cabannes observed. Ramm found only the triplet, no indication of any series of lines such as had been reported by E. Gross 470 and thought by him to be required by P. Debye's theory.⁴⁷¹ Ramm's conclusions rest on his study of the radiation that is scattered backward (turned 180° with reference to the incident light). Cabannes 467 has reported that the continuous background is almost completely depolarized.

In addition to those already mentioned, the theory and interpretation of such molecular scattering have been discussed by Y. Rocard, A. Bogros and Y. Rocard, J. Cabannes and Y. Rocard, W. Ramm, and A. Rousset,

Raman Scattering.—In the preceding cases, the spectrum of the scattered light was determined either solely by the medium or solely by the incident light as modified by the color of the scattering particles. But C. V. Raman ⁴⁷⁶ observed that the spectrum of the scattered light contains lines that are foreign to the spectrum both of the incident light and of the medium. The scattering that gives rise to these lines is known as the Raman scattering—sometimes in Germany as the Smekal-Raman scattering, A. Smekal ⁴⁷⁷ having shown theoretically in a letter on another subject, that a scattering of this type is demanded by the quantum theory. See also Y. Rocard.⁴⁷⁸

On the quantum theory, which accounts fairly well for the observed phenomena, when a quantum of radiation of frequency c_{v_i} strikes an atom

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406 Cabannes, J., and Salvaire, P., Compt. rend., 188, 907-908 (1929).
407 Cabannes, J., Idem, 191, 1123-1125 (1930).
408 Ramm, W., Physik. Z., 35, 756-773 (1934).
408 Brillouin, L., Ann. de Phys. (9), 17, 88-122 (1922).
470 Gross, E., Naturwissenschaften, 18, 718 (L) (1930).
471 Debye, P., Ann. d. Physik (4), 39, 789-839 (1912).
472 Rocard, Y., Ann. de Phys. (10), 10, 116-179, 181-231, 472-488 (1928).
473 Bogros, A., and Rocard, Y., Jour. de Phys. (6), 10, 72-77 (1929).
474 Cabannes, J., and Rocard, Y., Idem, 10, 52-71 (1929).
475 Rousset, A., Jour. de Phys. (7), 6, 507-515 (1935); Ann. de Phys. (11), 5, 5-135 (1936).
476 Raman, C. V., Indian J. Phys., 2, 387-398 (1928).
477 Smekal, A., Naturwissenschaften, 11, 873 875 (L) (1923).
478 Rocard, Y., Compt. rend., 186, 1107-1109 (1928).
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or an aggregation of atoms, there may be a transfer of a quantum of radiation of frequency c_{V_C} characteristic of the atom or aggregation. The transfer may occur in either direction, and c_{V_0} may be any of the characteristic frequencies of the material. Hence the scattered radiation will contain the additional frequencies $c(\nu_i + \nu_c)$ and $c(\nu_i - \nu_c)$, there being as many ν_c 's as there are frequencies characteristic of the material, and as many v_i 's as there are frequencies in the incident radiation. The complete set of additional frequencies attendant upon any one incident frequency (cv_i) is characterized by a definite set of frequency differences $(c\delta v)$, the same for every cvi. This set of frequency differences is called the Raman spectrum of the medium. It consists of two parts. In one, called the antistokes Raman spectrum, the actual frequencies exceed c_{ν_i} ; in the other, the stokes Raman spectrum, they are smaller than c_{ν_i} . Only those atoms and aggregations that are suitably excited can contribute to the first; whereas any that are unexcited may contribute to the second. Hence the stokes frequencies will be much the stronger, unless the medium is subjected to some definitely exciting action. Throughout the preceding, c = velocity of light, $\nu (\equiv 1/\lambda) = \text{wave-number}, \ \lambda = \text{wave-length of the radiation correspond-}$ ing to the frequency cv.

The relative intensities of the lines in either part—stokes or antistokes—of the Raman spectrum vary widely. Those corresponding to the fundamental frequencies of the medium are, in general, much stronger than those corresponding to the combination frequencies. On account of their low intensities, the Raman lines can be satisfactorily observed only in the laterally scattered radiation; they may be only partially plane-polarized, the amount of polarization varying from line to line. The appearance of a line, and its position, may vary with the temperature, and be changed by the addition of a solute.

Although the quantum theory accounts fairly well for the observed phenomena, it does not enable one to predict the Raman spectrum with certainty. The observed frequency differences seldom coincide exactly with the frequencies in the absorption spectrum; some are not represented in the absorption spectrum, and some of the frequencies found in that are not represented in the Raman spectrum. Nevertheless, the Raman effect enables one to obtain from studies in the visible and in the near ultraviolet spectrum much information regarding those characteristic vibrations that would otherwise have to be studied in the infrared, where observations are much more difficult. Therein lies one reason for the importance that is attached to the study of Raman spectra. Bibliographies are published from time to time in the *Indian Journal of Physics*, and the status of the entire subject in 1931 has been set forth and discussed by K. W. F. Kohlrausch.

⁴⁷⁹ See also, Raman, C. V., *Indian J. Phys.*, **6**, 263-273 (1931); Raman, C. V., and Krishnan, K. S., *Idem*, **2**, 399-419 (1928); Ganesan, A. S., and Venkateswaran, S., *Idem*, **4**, 195-280 (1929); Bhagavantam, S., *Idem*, **5**, 237-307 (1930).

⁴⁵⁰ Kohlrausch, K. W. F., "Der Smekal-Raman Effekt," Springer, Berlin, 1931.

Electron and β-ray Luminescence.—See page 317. Mechanical Luminescence.—See page 317.

Fluorescence of Water.

When water is subjected to gamma-rays from radium, it emits a white luminescence that is visible to the dark-adapted eye, and that is more strongly absorbed by 1 mm of glass than by 5 mm of either quartz or rock salt. The spectrum of this luminescence is continuous throughout the range covered by observations (visible spectrum and ultraviolet to $\lambda=2500\mathrm{A}$), and probably extends to the shortest wave-length unabsorbed by water. It is richer in the short waves than is the radiation from a 0.5-watt incandescent electric lamp.⁴⁸¹

P. A. Čerenkov (also spelled Tscherenkow)⁴⁸² has reported (1934) that the luminescence excited in water by γ -rays is partially polarized, the electric vector lying parallel to the line of propagation of the incident radiation, and is not reduced by the common quenchers of fluorescence (KI, AgNO₃, nitrobenzene), nor by heating to 100 °C. He has reported (1936) effects produced by a strong magnetic field, and has concluded (1937) that the luminescence arises from the action of the Compton electrons freed by the γ -rays, as was suggested by S. Wawilow,⁴⁸³ and not by the γ -rays themselves. See also p. 317.

Irradiating water by x-rays gave rise to no luminescence (Čerenkov; 1934).

- In 1925, K. S. Krishnan ⁴⁸⁴ concluded that when a beam of light is passed through water the laterally scattered light contains fluorescent radiation; and Y. Rocard ⁴⁸⁵ came to the same conclusion. The latter decided that this fluorescence is not due to $(H_2O)_n$ molecules, but probably to the presence of glass dissolved from the container.
- S. J. Wawilow and L. A. Tunumermann ⁴⁸⁶ have found that the fluorescence of water, which they describe as blue, is reduced very little if at all by boiling, but repeated distillation in quartz completely destroys it. Bubbling of either air or CO₂ through doubly distilled water increases the intensity of the fluorescence, but the bubbling of oxygen does not. This "fluorescence" may include the scattered light also.
- A. Carrelli, P. Pringsheim, and B. Rosen ⁴⁸⁷ have stated that Berlin city water excited by $\lambda = 3650 \text{A}$ exhibits a rather strong blue-violet fluorescence consisting of a very broad, ill-defined, continuous spectral band. The same fluorescence of essentially the same intensity was obtained with Kahlbaum's distilled water, but was almost absent from his conductivity water.

 ⁴⁸¹ Mallet, L., Compt. rend., 183, 274-275 (1926); 187, 222-223 (1928); 188, 445-447 (1929).
 482 Čerenkov, P. A. (also spelled Tscherenkow), Compt. rend. Acad. Sci. URSS (N. S.), 1934, 455-457 (1934); 12, 413-416 (1936); 14, 101-105 (1937).

¹⁵³ Wawilow, S., Idem, 1934, 459 461 (1934).

⁴⁸⁴ Krishnan, K. S., Phil. Mag (6), 50, 697-715 (1925).

⁴⁵ Rocard, Y., Compt. rend., 180, 52-53 (1925).

⁴⁹⁰ Wawilow, S. J., and Tummermann, L. A., Z. Physik, 54, 270-276 (1929).

⁴⁸⁷ Carrelli, A., Pringsheim, P., and Rosen, B., Z. Physik, 51, 511-519 (1928).

The interpretation of observations purporting to measure the amount of fluorescence excited in water by the optical spectrum is difficult. The radiation scattered by the Tyndall, Rayleigh, and Raman effects being incompletely polarized, the presence of an unpolarized component in the scattered light is not a certain criterion for even the presence of fluorescence. if the depolarization factor ρ is measured once for the total laterally emitted light, and again for the same light deprived solely of the fluorescent light, then from these two factors the ratio of the intensity (2f) of the fluorescent light to that (a + b) of the scattered can be determined, a and b being the I_w and the I_w of the scattered light (p. 300). In the first case, $\rho' =$ (a+f)/(b+f); and in the second, $\rho = a/b$; whence 2f/(a+b) = $2(\rho'-\rho)/(1-\rho')\cdot(1+\rho)$. In practice, the fluorescent light is removed by means of a filter cutting out the ultraviolet. When the filter is in the incident beam, ρ is measured; when in the scattered, ρ' . This assumes that the intensities of the additional Raman bands that are present in the second case contribute negligibly to the intensity of the scattered light. By this procedure the following results were obtained:

2f/(a+b)	Filter	Reference
0.069	Orange	Krishnan, K. S. 484
0.033	Green	Ibid.
0.03	Quinine	Sweitzer, C. W., J. Phys'l Chem., 31, 1150-1191 (1927).
0.13	Quinine(?)	Canals, E., and Peyrot, F., Compt. rend., 198, 1992-1994 (1934).

Rayleigh Scattering by Water.

In obtaining the data given in the following tables, the investigators made no attempt to eliminate the effects of the Raman scattering or of fluorescence, except as is indicated; and the data for polarization likewise refer to the total laterally emitted light.

The intensity of the scattered light varies reversibly with the temperature, decreasing as the temperature rises, 488 but the published data cannot be interpreted quantitatively.

E. O. Hulburt ⁴⁸⁰ has shown that the observations by W. Beebe and G. Hollister ⁴⁹⁰ of the intensity of the light scattered horizontally by the sea at various depths can be satisfactorily accounted for by the Rayleigh scattering and the values he himself obtained for the coefficient of absorption, except in the first 250 feet, where the absorption has to be increased by an amount equivalent to the presence in each cubic centimeter of one mote one-tenth of a square millimeter in sectional area.

More recently, L. H. Dawson and E. O. Hulburt ⁴⁹¹ have found that within their experimental error (< 15%) the total light scattered by water in the range $\lambda = 2536 \text{A}$ to 5790A varies with λ as demanded by the Einstein-Smoluchowski expression (see Table 148).

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<sup>488</sup> Schade, H., and Lohfert, H., Kolloid Z., 51, 65-71 (1930).

<sup>489</sup> Hulburt, E. O., J. Opt. Soc. Amer., 22, 408-417 (1932).
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⁴⁸⁰ Beebe, W., and Hollister, G., Bull. N. Y. Zool. Soc., 33, 249-263 (1930).

⁶⁰¹ Dawson, L. H., and Hulburt, E. O., J. Opt. Soc. Amer., 27, 199-201 (1937) → Phys. Rev. (2), 51, 1017 (A) (1937).

In 1930, J. Plotnikow and L. Splait 492 described and studied what they called a longitudinal scattering. The work was continued by J. Plotnikow and S. Nishigishi 493 and others, with varying results, leading to a discussion, sometimes spirited. The original contention of Plotnikow and his associates has been upheld by B. Čoban 494 and by L. Šplait, 494a but it seems almost certain that everything that is not spurious in the phenomena described can be accounted for by the well-known Tyndall and Rayleigh scattering.495

Table 148.—Polarization and Intensity of Light Laterally Scattered by Water

The values credited to K and to RR (see refer-See remarks in text. ences^a) are essentially those given in the compilation by J. W. T. Walsh and H. Buckley. 496

I = intensity of the laterally scattered light at a distance r from thevolume V of water from which the light comes; E = intensity of the incident (exciting) light; $\rho =$ depolarization factor.

 I_w/I_f = ratio of intensity of light laterally scattered by water to that of the light similarly scattered by the indicated fluid under the same conditions; l = liquid, g = gas at 0 °C and 1 atm.

 ρ_i , ρ_s = value of ρ when the indicated filter is in the incident, scattered, beam, respectively. The filter is specified by the color of the light transmitted.

 λ = wave-length of the incident (exciting) light.

Unit of $\lambda = 1$ A = 10^{-8} cm; ρ is a pure number.

I. Polarization

T. Toldi	Eution.				
Unfilte	red		Filt	ered -	
ρ	Ref.	Ref.a→	G	K	K
		Filter	ρ_{i}	$\rho_{\mathbf{i}}$	Pa
0.067	\mathbf{M}^{1}	Red	0.119		
0.05	C	Orange	0.108	0.085	0.118
0.125	RR	Green	0.105	0.079	0.095
0.106	G	Blue-gr.	0.105	****	
0.096	$\check{\mathbf{K}}$	Blue	0.144	0.145	0.099
0.116	Ro	2.40	0.111	******	
0.109	S				
0.096	Ra				
II. Inten	isity.				
Fluid	I_w/I_f	Ref.a	λ	$10^6 Ir^2/VE$	Ref.a
Ether (l)	0.192	ML		1.77	ML
Ether (l)	0.192	RR	4358	1.77	M^2
Benzene (l)	0.069	S	5461	0.72	M^2
Toluene (l)	0.060	\mathbf{M}^{1}	5780	0.57	M^2
	165.	RR	5700	0.01	
Air (g)	100.	1/1/			

⁴⁸² Plotnikow, J., and Splait, L., Physik. Z., 31, 369-372 (1930).

⁴⁹⁸ Plotnikow, J., and Nishigishi, S., Idem, 32, 434-444 (1931).

⁴⁸⁴ Coban, B., Acta Phys. Polon., 4, 1-16 (1935).

⁴⁹⁴a Splait, L., Idem, 4, 329-330 (1935). Krishnan, R. S., Proc. Indian Acad. Sci., 1, 44-47, 211-216 (1935); Mitra, S. M., Z. Physik,
36, 34-36 (1935); Vrkljan, V. S., Acta Phys. Polon., 4, 325-327 (1935); and Katalinić, M., Koll. Z.,
28-296 (1936); Z. Physik, 106, 439-452 (1937).
Walsh, J. W. T., and Buckley, H., Int. Crit. Tables, 5, 266 (1929).

Table 148-(Continued)

III. L. H. Dawson and E. O. Hulburt.⁴⁹¹ If the molecule of water were isotropic, then, on the density-fluctuation theory of Einstein and Smoluchowski, the total radiation of wave-length λ that a unit volume of water at 22 °C would scatter per unit of solid angle (i.e., per steradian) in directions perpendicular to the direction of propagation of the incident light would be $I_{\lambda}\alpha$, where α has the values here tabulated, and I_{λ} is the intensity of the incident radiation of wave-length λ . Since the molecules are anisotropic, the scattering will exceed $I_{\lambda}\alpha$. The amount of this excess in the ultraviolet is unknown; in the visible spectrum it is less than 30 per cent. The observations of Dawson and Hulburt in the range $\lambda = 253.6$ to 546.1 m μ agree relatively with the values here tabulated.

	Unit of $\lambda =$	$1 \text{ m}\mu = 10^{-7} \text{ c}$	m ; of $\alpha = 1$ cm	-1 steradian-1.	Temp. = 22	°C = 295 °K	
λ	$10^7\alpha$	λ	107a	λ	$10^7\alpha$	λ	107a
600	0.79	500	1.66	400	4.06	300	14.8
550	1.03	450	2.53	350	7.15	250	34.4
" Refer	ences:						
С	Cabannes	. J., Jour.	le Phys. (6).	3, 429-442 (1922).		
C G				(1923) ← Coi		e cienc. Uniz	. La Plata
	(Mat. Fi	is.), 3, 251-3	15 (1923); Z	. Physik, 30,	231-239 (19.	24).	
K	Krishnan	, K. S., Phil	. Mag. (6),	50, 697-715 (1	1925).		
\mathbf{M}	Martin,	W. H., M1=	J. Phys'l Cl	hem., 24, 478.	492 (1920)	M2-Idem.	26, 471-476
	(1922).	•			• •		
ML	Martin, V	W. H., and I	Lehrman, S.,	Idem, 26, 75-1	88 (1922).		
Ra				. 4, 15-38 (19			
D.,			J 190 F) F				

Ro Rocard, Y., Compt. rend., 180, 52-53 (1925).

RR Raman, C. V., and Rao, K. S., Phil. Mag. (6), 45, 625-640 (1923).

S Sweitzer, C. W., J. Phys'l Chem., 31, 1150-1191 (1927).

Rocard reports this value for green light, and says that ρ varies very little with the wave-length. If fluorescence had not been eliminated [?] then $\rho = 0.16$ to 0.18.

Raman Scattering by Water.

Water contrasts sharply with most other liquids in that its Raman spectrum consists of broad diffuse bands, some of which overlap. This, together with the fact that the spectrum of the mercury arc, which is the most satisfactory illuminant, contains a number of bright lines, makes interpretation of the observations difficult, unless care is taken to remove from the light of the arc all except one line, or a small group of closely spaced lines. That has been done by H. Hulubei, 497 H. Hulubei and Y. Cauchois, 498 M. Magat, 499 and J. H. Hibben 500; but most of the recorded observations have been made with the unfiltered radiation.

The most prominent features of the Raman spectrum of water are two bands, one broad and centered near the wave-length corresponding to $\delta\nu=3400~{\rm cm^{-1}}$, the other narrow and centered near $\delta\nu=1650~{\rm cm^{-1}}$. There has been much discussion about the structure of the first (see Table 152). It probably has three components, the strongest having its maximum near $\delta\nu=3400~{\rm cm^{-1}}$, the one of intermediate strength near $\delta\nu=3200~{\rm cm^{-1}}$, and

⁴⁹⁷ Hulubei, H., Compt. rend., 194, 1474-1477 (1932).

⁴⁰⁸ Hulubei, H., and Cauchois, Y., Idem, 192, 1640-1643 (1931).

⁴⁰⁰ Magat, M., Idem, 196, 1981-1983 (1933); Jour. de Phys. (7), 5, 347-356 (1934).

⁸⁰⁰ Hibben, J. H., J. Chem'l Phys., 5, 166-172, 994 (1937).

a very weak one near $\delta_{\nu}=3600~{\rm cm^{-1}}$, but H. Hulubei 407 and M. Magat 501 failed to find the 3600 cm⁻¹ component, although they sought for it; and E. H. L. Meyer 502 suggested that the apparent structure of this band is an optical illusion. For the early discussion of the subject see W. Gerlach 503 and E. H. L. Meyer. 504 More recently, Magat 505 has reported that he has found this 3600 cm⁻¹ component, but only at temperatures above 37 °C.

I. R. Rao ⁵⁰⁶ has sought to interpret the observations in terms of the polymerization of water; M. Magat, ⁵⁰¹, ⁵⁰⁵, ⁵⁰⁷ accepting the quasicrystal-line theory of liquid structure, has sought to interpret them in terms of the several modes of vibration of the molecule when subjected to the action of its neighbors. Magat's view is the one more favored at present.

Most of the early work was limited to a study of the bands near $\delta_{\nu}=3400$ and 1650 cm⁻¹, but many other lines and bands have been mapped (see Table 153). These have been regarded by I. R. Rao and P. Koteswaram ⁵⁰⁸ as spurious, as arising from excitation by another spectral line than that supposed by the observer; but J. H. Hibben ⁵⁰⁹ seems to have shown conclusively that such is not the case, that in his work, at least, the lines in dispute cannot have arisen from excitation by any other line than that he supposed, and that at least the lines near $\delta_{\nu}=175$, 500, 1659, and 2150 cm⁻¹ are true Raman lines. He had not observed the lines reported at $\delta_{\nu}=4023$ and 5100 cm⁻¹, but he gave reasons for believing that Rao and Koteswaram's criticism is inapplicable to them also.

G. Bolla 510 has reported a spurious multiplication of the Raman bands under certain instrumental conditions.

General reviews of certain phases of the work in this field have been recently published by M. Magat,⁵¹¹ A. Kastler,⁵¹² P. C. Cross, J. Burnham, and P. A. Leighton,⁵¹³ and J. H. Hibben.⁵¹⁴

Polarization and Intensity of the Raman Bands of Water.—J. Cabannes 515 has found that for a given substance the polarization of any given Raman line or band is independent of the frequency of the exciting radiation, and that the amount of the polarization of the several Raman lines or bands, each corresponding to a different value of δ_{ν} , may differ, the depolarization factor (ρ) lying between 0 and 1 (actually, 6/7 is the limiting

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**Bold Magat, M., Jour. de Phys. (7), 5, 347-356 (1934).

**Board Meyer, E. H. L., Physik. Z., 31, 510-511 (1930).

**Board Meyer, E. H. L., Idem, 31, 699-698 (1930).

**Board Magat, M., Jour. de Phys. (7), 6, 648-65S (1935).

**Boord Rao, I. R., Proc. Roy. Soc. (London) (A), 130, 489-499 (1931); Nature, 132, 480 (1933);

**Proc. Roy. Soc. (London) (A), 145, 489-508 (1934); Phil. Mag. (7), 17, 1113-1134 (1934).

**Boord Magat, M., Ann. de Phys. (11), 6, 108-193 (1936); Trans. Faraday Soc., 33, 114-120 (1937).

**Boord Hibben, J. H., Idem, 5, 994 (L) (1937).

**Boord Hibben, J. H., Idem, 5, 994 (L) (1937).

**Boord Hibben, J. H., Idem, 5, 994 (L) (1931); 129, 60 (L) (1932).

**Bill Magat, M., Ann. de Phys. (11), 6, 108-193 (1936) (Bibliog. of 148 entries).

**Bill Kastler, A., Rev. gén. dcs. Sci. (Paris), 47, 522-536, 559-566 (1936).

**Bill Cross, P. C., Burnham, J., and Leighton, P. A., J. Am. Chem. Soc., 59, 1134-1147 (1937).

**Bill Cabannes, J., Compt. rend., 187, 654-656 (1918).
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value). Values of ρ are given in Table 149. See also F. Heidenreich.⁵¹⁶ Using $\lambda_{Hg} = 3650$ A as exciter, G. I. Pokrowski and E. A. Gordon 517 measured both the polarization $(1 - \rho)$ and the relative intensity of the band $\delta_{\nu} = 3400 \text{ cm}^{-1}$ when scattered at an angle θ with the direction of the incident beam (Table 149).

Of the three components ($\delta_{\nu} = 3200, 3450, 3600 \text{ cm}^{-1}$) of the $\delta_{\nu} = 3400$ cm⁻¹ band, the second is the strongest at room temperatures; and the third the weakest.518

Table 149.—Polarization and Angular Scattering of the Raman Bands of Water

 $\delta v = \text{approximate shift defining the maximum of the band or component}$ studied; ρ = depolarizing factor; I = relative intensity of the band as scattered at an angle θ with the direction of the incident beam.

		Uni	t of $\delta \nu = 1$ cm	n-1; p and I are	ratios		
δν→ Ref.α	175	500	750	1650	3200	3450	3600
An	0.85	g	g	_	0.10-0.15	0.40-0.50	0.85
CdeR	-	-	_	0.4	$< 0.30^{b}$	0.30	_
Ra	-	-		_	0.60	0.48	0.75
Sp CR	-		-		0.62	0.52	0.54
CR	-	-	-	-	Yese	Noc	-
PGª Exc	iter, λπg =	= 3650A; δν	= 3400.				
θ	20°	40°	60°	90°	120°	140°	160°
I	2.0	1.9	1.3	1.0	0.7	0.7	0.7
$(1-\mu)^d$	0.07	-	_	0.70 ± 0.02		_	0.7

a References :

An Ananthakrishnan, R., Proc. Indian Acad. Sci., 3, 201-205 (1936).

(Der Cabannes, J., and de Riols, J., Compt. rend., 198, 30-32 (1934).

(Cabannes, J., and Rousset, A., Idem, 194, 706-708 (1932).

PG Pokrowski, G. I., and Gordon, E. A., Ann. d. Physik (5), 4, 488-492 (1930).

Ra Ramaswamy, C⁵¹⁸

Sp Specchia, O., Nuovo Cim. (N. S.), 9, 133-137 (1932).

This line is said to be more polarized than the 3450 cm⁻¹ one, hence the < 0.30.

No numerical value given; the 3200 line is said to be depolarized; the 3450 one

They call these values the "polarization," presumably meaning $(1 - \rho)$.

Effect of Temperature on the Raman Scattering by Water.—At 11.5 °C the intensity of the band $\delta v = 3400 \text{ cm}^{-1}$ is the same whether the band is excited by $\lambda_{Hg} = 3020$, 2968, or 2654A; whereas at 55 °C the one excited by $\lambda_{\rm Hg} = 2968$ A is about 20 per cent more intense than either of the others.⁵¹⁹ But P. Pringsheim and S. Slivitch 520 have reported that the relative intensities of the several repetitions of a given band ($\delta v = 3400 \text{ cm}^{-1}$), each corresponding to one of the stronger lines of the mercury spectrum, are independent of the temperature of the water.

Both of the two broad bands observed at 4690A and 4250A (presumably

517 Pokrowski, G. I., and Gordon, E. A., Ann. d. Physik (5), 4, 488-492 (1930).

⁵¹⁰ Heidenreich, F., Z. Physik, 97, 277-299 (1935).

⁵¹⁸ Ramaswamy, C., Nature, 127, 558 (L) (1931); Specchia, O., Nuovo Cim. (N. S.), 9, 133-137 (1932). See also Hibben, J. H., J. Chem'l Phys., 5, 166-172 (1937).

⁵¹⁰ Meyer, E. H. L., and Port, I., Physik. Z., 31, 509-510 (1930).

⁸²⁰ Pringsheim, P., and Slivitch, S., Z. Physik, 60, 581-585 (1930).

corresponding, respectively, to δν about 3400 cm⁻¹ and 1650 cm⁻¹) become narrower and sharper as the temperature is increased.⁵²¹

The $\delta v = 140 \text{ cm}^{-1}$ band, observed and studied between 4 °C and 97 °C by E. Segrè, 522 decreases in intensity as the temperature is increased.

As the temperature is increased, the maximum of the broad Raman band centered near $\delta_{\nu} = 3400$ shifts in the direction of increasing δ_{ν} , and

Table 150.—Shift of Raman Lines of Water with Change in Temperature

(See text for comments and references to other work.)

cδν is the frequency difference corresponding to the maximum intensity of the Raman line or band; $\delta_{\nu} = \pm (1/\lambda_m - 1/\lambda_i)$, where λ_i and λ_m are the wave-lengths of the incident and the scattered radiation, respectively; c = velocity of light; $\rho = \text{density of the water}$.

Unit of $\delta \nu =$	= 1 cm ⁻¹ ; or ρ = 1 g/ml.	$lemp. = l^{-1}C$		
U	k ^b		_ Raa _	
ρ	δν	t		δν
0.98	3448	0		3502
0.93	3497	4		3412
0.86	3524	38		3493
0.78	3520	98		3466
0.70	3530			
0.66	3528		Spa _	
	3530	17		3406
0.33	3530	41		3417
0.133	3530	60		3429
	(3530°	80		3452
0.096	3646	91		3474
0.055	3646^{d}			
0.025	3645		- Mea -	
	(3639¢	11.5		3414
0.0135	13653	55		3430
0.007	(3639*	92		3551
0.007	3653			
	0.98 0.93 0.86 0.78 0.70 0.66 	0.98 3448 0.93 3448 0.93 3497 0.86 3524 0.78 3520 0.70 3530 0.66 3528 - 3530 0.133 3530 0.133 3530 0.096 {3530° 3646 0.055 3646 ^d 0.025 3645 0.0135 {3639° 3639°	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.98 3448 0 0.93 3497 4 0.86 3524 38 0.78 3520 98 0.70 3530 0.66 3528

	H	ia	
28 °C	88 °C	28 °C	88 °C
144	149	2170	2118
440	450	3219	3222
1627	1629	3445	3460

a References:

Hi

Me

Ra

Hibben, I. H.⁵⁰⁰
Meyer, E. H. L., Physik. Z., 31, 510-511 (1930).
Rao, I. R., Proc. Roy. Soc. (London) (A), 145, 489-508 (1934).
Specchia, O., Nuovo Cim. (N. S.), 7, 388-391 (1930).
Ukholin, S. A., Compt. rend. Acad. Sci. URSS, 16, 395-398 (1937).

b Ukholin worked with water sealed in quartz tubes and heated to various temperatures; the state of the water was specified by means of the temperature and the density, which in many cases was less than at the critical point ($\rho_{crit} = 0.33$).

[&]quot;Here the band persists and a new line appears.

⁴ Here the band has vanished and only the new line remains.

The line has now split into two.

⁵²¹ Meyer, E. H. L., Physik. Z., 30, 170 (1929).

⁵²² Segrè, E., Atti Accad. Linc. (6), 13, 929-931 (1931).

the component of smallest δ_{ν} (δ_{ν} about 3200 cm⁻¹, corresponding to $\lambda_{R}=3.13~\mu$) decreases in intensity, nearly vanishing as 100 °C is approached.⁵²³ The intensities of the other two components of that band (δ_{ν} about 3450 cm⁻¹ and 3600 cm⁻¹, corresponding to $\lambda_{R}=2.90~\mu$ and 2.77 μ , respectively) change but little with the temperature,⁵²⁴ the change of the last ($\delta_{\nu}=3600~{\rm cm^{-1}}$) being an increase (Rao 1930).⁵²³

The original papers should be consulted. More recent work, covering the effect of temperature on each of several bands, may be found in the papers here noted.⁵²⁵

Certain data given by Hibben and Ukholin are included in Table 150. Magat ⁵²⁶ is of the opinion that the variation of the Raman lines of water with the temperature is peculiar near 40 °C; but G. Bolla ⁵²⁵ disagrees with him.

Effect of Solutes on the Raman Scattering by Water.—The relative intensities of the several repetitions of a given band ($\delta \nu = 3400~{\rm cm}^{-1}$), each corresponding to one of the stronger lines of the mercury spectrum, are unchanged by the addition of a solute to the water.⁵²⁰

The solution in water of HNO₃ or of certain salts forming electrolytic solutions causes the components of the 3400 cm⁻¹ band to become sharper, the 3200 cm⁻¹ component to decrease in intensity, and the 3600 cm⁻¹ component to increase; the band is shifted in the direction of increasing δ_{ν} . In concentrated solutions of HNO₃, the 3200 cm⁻¹ component is vanishingly weak; whereas the 3600 cm⁻¹ one is the strongest of the three.⁵²⁷ In solutions of NaNO₃, this band is shifted as just stated, but the intensity of the 3600 cm⁻¹ component is decreased, vanishing in a 66 per cent solution.⁵²⁸

On the other hand, the solution of HCl decreases the intensity of both the components 3200 cm⁻¹ and 3600 cm⁻¹, and somewhat increases that of 3400 cm⁻¹ (Brunetti and Ollano ⁵²⁷; Rafalowski ⁵²⁷).

In contrast to the observers mentioned in the two preceding paragraphs, W. Gerlach,⁵³⁰ who has reported that he finds only two components (3200 cm⁻¹, 3400 cm⁻¹) in the 3400 cm⁻¹ band, has stated that only the 3400 cm⁻¹ component was visible in solutions of LiCl and of CaCl₂, but both were visible in solutions of ZnCl₂ and of CdCl₂. In solutions of the alkali nitrates

Bhagavantam, S., Ind. J. Phys., 5, 49-57 (1930); Meyer, E. H. L., Physik. Z., 31, 510-511 (1930); Nisi, H., Jap. J. Phys., 7, 1-32 (1931); Rao, I. R., Nature, 125, 600 (1930); Proc. Roy. Soc. (London) (A), 130, 489-499 (1931); Specchia, O., Nuovo Cim. (N. S.), 7, 388-391 (1930); Ganesan, A. S., and Venkateswaran, S., Indian J. Phys., 4, 195-280 (1929).

Boulesan, A. S., and venkaueswaran, S., Indian J. Phys., 4, 195-280 (1929).

Bold Bhagavantam, S., Indian J. Phys. 5, 49-57 (1930); Ganesan, A. S., and Venkateswaran, S., idem, 4, 195-280 (1929).

Bolla, G., Nuovo Cim. (N. S.), 12, 243-246 (1935); Rao, C. S. S., Proc. Roy. Soc. (London) (A), 151, 167-178 (1935); Magat, M., Jour. de Phys. (7), 6, 648-65S (1935); Ann. de Phys. (11), 6, 108-193 (1936); Ananthakrishnan, R., Proc. Indian Acad. Sci., 3, 201-205 (1936); Cross, P. C., Burnham, J., and Leighton, P. A., J. Am. Chem. Soc., 59, 1134-1147 (1937); Hibben, J. H., J. Chem'l Phys., 5, 166-172 (1937); Magat, M., Trans. Faraday Soc., 33, 114-120 (1937); Ukholin, S. A., Compt. rend. Acad. Sci. URSS, 16, 395-398 (1937).

⁸³⁰ Magat, M., Jour. de Phys. (7), 6, 64S-65S (1935); Trans. Faraday Soc., 33, 114-120 (1937).
837 Brunetti, R., and Ollano, Z., Atti Accad. Lincci (6), 12, 522-529 (1930); Rafalowski, S.,
Bull. Int. dc l'Acad. Polonaise (A), 1931, 623-628 (1931) → Nature, 128, 546 (1931); Rao, I. R.,
Nature, 124, 762 (1929) ← Proc. Roy. Soc. (London) (A), 127, 279-289 (1930); Nature, 125, 600 (1930); Proc. Roy. Soc. (London) (A), 130, 489-499 (1931).

⁸⁰⁸ Cahannes, J., and de Riols, J., Compt. rend., 198, 30-32 (1934).

Reo Gerlach, W., Naturwissenschaften, 18, 68 (L), 182-183 (L) (1930); Physik. Z., 31, 695-698 (1930).

the wave-length separation of the two components increased almost linearly with the concentration, the band shifted in the direction of decreasing δ_{ν} , and the 3400 cm⁻¹ component split into two, as the concentration increased. He stated that a broad unresolved band at $\lambda = 4160 \text{A}$ (Is it the $\delta_{\nu} = 1650 \text{ cm}^{-1}$ band?) is weak in solutions of LiCl, but is sharp and displaced in the direction of increasing δ_{ν} in solutions of CaCl₂. N. Embirikos ⁵³¹ generally confirms Gerlach.

In ammonium solutions, the $\delta_{\nu}=1650~\rm cm^{-1}$ band is shifted in the direction of increasing δ_{ν} , the shift being small for the nitrate and the chloride, but exceptionally great (about 30 cm⁻¹) for the sulfate.⁵³² This band is but little affected by adding NaNO₃ to the water.⁵²⁸

Table 151.—Analysis of the Raman Spectrum of Water 542

(For the band near $\delta v = 3400$ cm ¹ see Table 152.)

 v_{σ} , v_{π} , and v_{δ} are the quantum numbers corresponding to the three fundamental modes of vibration of the free molecule (Table 64); v_1 , v_2 , v_3 , and v_0 are four others corresponding to fundamental vibrations determined by the interaction of the molecule with its neighbors. The frequency of a given vibration is c_{ν} where $\nu = v_{\sigma}v_{\sigma} + v_{\pi}v_{\pi} + \ldots + v_{0}v_{0}$, and $c_{\nu}v_{\sigma}$, $c_{\nu}v_{\pi}$, $c_{\nu}v_{0}$ are the frequencies of the 7 fundamental vibrations.

ve	U#	vs	v_1	v ₂	v	vo	Infrared	Raman	
							- νυ	las ———	Peale
0	0	0	0	1	0	0		60	(60)
0	0	0	0	0	0	1	-	152-225	166a
0	0	0	0	0	1	0	510	500	570ª
Ŏ	Ō	0	i	0	0	0	670	740	(700)
Õ	Ö	1	ō	1	Ö	0	1710		1720
0	0	1	0	0	0	1	1850	*****	1820
Ō	0	1	Ô	0	1	0	2135	2135	2160
ĭ	Õ	Ō	Ŏ	Ö	Ĩ	Õ	4023	4024	3950
ĩ	ŏ	1	ŏ	ő	î	Ö	5590		5620

* Frequencies near these may be derived from Bernal and Fowler's proposed structure of water (see p. 174).

See also C. C. Hatley and D. Callihan,⁵³³ H. Hulubei,⁴⁹⁷ C. Ramaswamy,⁵³⁴ A. da Silveira and E. Bauer,⁵³⁵ A. Hollaender and J. W. Williams,⁵³⁶ E. Bauer,⁵³⁷ M. Magat,⁵³⁸ F. Cennamo,⁵³⁰ J. H. Hibben,⁵⁰⁰ P. A. Leighton and J. Burnham,⁵⁴⁰ and for the earlier work, especially K. W. F. Kohlrausch.⁴⁸⁰

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531 Embirikos, N., Physik. Z., 33, 946-947 (1932).
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⁵⁸² da Silveira, A., Compt. rend., 195, 521-523 (1932).

⁵⁸⁸ Hatley, C. C., and Callihan, D., Phys. Rev. (2), 38, 909-913 (1931).

⁸⁸⁴ Ramaswamy, C., Indian J. Phys., 5, 193-206 (1930).

tat da Silveira, A., and Bauer, E., Compt. rend., 195, 416-418 (1932).

⁵⁸⁶ Hollaender, A., and Williams, J. W., Phys. Rev. (2), 34, 994-996 (1929).

⁸⁸⁷ Bauer, E., Jour. de Phys. (7), 6, 63S-64S (1935).

⁶⁹⁸ Magat, M., Jour. de Phys. (7), 6, 64S-65S (1935).

⁵³⁹ Cennamo, F., Nuovo Cim. (N. S.), 13, 304-309 (1936).

⁵⁴⁰ Leighton, P. A., and Burnham, J., J. Am. Chem. Soc., 59, 424-425 (L) (1937).

Interpretation of the Raman Spectrum of Water.—Numerous attempts have been made to interpret the Raman spectrum in terms of the fundamental vibrations of the water molecule (see Tables 64 and 65) and in terms of the observed infrared spectrum of water. At first they were limited to the band near $\delta v = 3400 \text{ cm}^{-1}$ (see Table 152), but recently they have been extended by M. Magat 541 to other lines. He has concluded that

Table 152.—Analysis of the Raman Band near $\delta \nu = 3400$ cm⁻¹

The following values of $\delta\nu$ and of $\lambda_R \equiv 1/\delta\nu$ are those assigned by the several observers to the maxima of the components of the band. Some give $\delta\nu$; some, λ_R ; and some, both; if only one of these quantities has been published, the compiler has computed the other from it. Such computed values are enclosed in parentheses. Hu and Mag sought for the 3600 component, but could not find it; the others cited in the second section of the table say nothing about it. E. H. L. Meyer ⁵⁴³ suggested that the apparent structure of this band is an optical illusion; for the resulting controversy, see W. Gerlach ⁵⁰³ and E. H. L. Meyer. ⁵⁰⁴

		Unit of $\delta \nu = 1$ c	m^{-1} ; of $\lambda_R = 1 \mu$	= 10 ⁻⁴ cm		
				· λ _R		Ref.
(3195)	(3448)	(3610)	3.13	2.90	2.77	$\mathbf{Bh^1}$
3200	3435	3630	3.12	2.91	2.75	Bo3,4
3225	3469	3589	3.10	2.88	2.79	BO
3224	3436	3625	(3.10)	(2.91)	(2.76)	CRi
3230	3450	3560	(3.10)	(2.90)	(2.81)	CC
3199	3453	3609	3.13	2.90	2.77	GaV
3228	3435	3624	3.10	2.91	2.76	HaC
3206	3456	3578	3.12	2.89	2.79	N_1
3180	3440	3630	(3.14)	(2.91)	(2.75)	$\mathbf{R}\mathbf{y^i}$
3208	3419	3582	(3.12)	(2.92)	(2.79)	Ros
(3205)	(3413)	(3584)	3.12	2.93	2.79	Ro4
3084	3423	3628	3.24	2.92	2.75	Ro^5
3217	3433	3582	(3.11)	(2.91)	(2.79)	Ros.7
3278	3406	3569	3.04	2.92	2.80	Sp¹ Sp¹
3246	3405	3554	3.08	2.93	2.81	Sp^1
3208	3435	3595	3.117	2.90,	2.77,	Mean
3232	3422	-	(3.09)	(2.92)		Ge
3324	3513		(3.01)	(2.85)	-	DuKo1,2
3233	3443	none	(3.09)	(2.90)		Hu
3195	3394		(3.13)	(2.95)		Ki
3221	3435	none	(3.10)	(2.91)		Mag ²
31956	3437		(3.13)	(2.91)	-	N^2
3217	3441		3.091	2.90,		Mean

^{*} References: See Table 153.

^b Nisi states that when the exciter is $\lambda_{Hg} = 4047$ A or 3650–3663A the band looks like a triplet, but that when the exciter is $\lambda_{Hg} = 2967$ A it consists of only two (diffuse) bands, as here given, with no indication of a third. Magat (Mag ^a) used both $\lambda_{Hg} = 4358$ A and $\lambda_{Hg} = 2537$ A, and in both cases failed to find the 3600 cm⁻¹ component.

⁸⁴¹ Magat, M., Ann. de Phys. (11), 6, 108-193 (1936); Trans. Faraday Soc., 33, 114-120 (1937).

⁵⁴⁸ Magat, M., Ann. de Phys. (11), 6, 108-193 (1936).

⁸⁴⁸ Meyer, E. H. L., Physik. Z., 31, 510-511 (1930).

in addition to the three independent frequencies of the free molecule, four others, representing the effects of other molecules upon the one that is scattering the radiation, must be considered. His scheme is given in Table 151. S. A. Ukholin 525 has concluded that it is unsatisfactory to ascribe the maxima near $\delta \nu = 3400$ and 3600 cm⁻¹, respectively, to Bernal and Fowler's types II and III of water (p. 174). See also R. Ananthakrishnan. 525

Table 153.—Raman Spectrum of Water

Here are given all the more important reported values of δ_{ν} and of $\lambda_{R} (\equiv 1/\delta_{\nu})$. Some refer to the maxima of unresolved bands, some to the maxima of the components of bands.

Unit of $\delta v = 1$ cm⁻¹: of $\lambda v = 1$ v = 0.0001 cm

			Un	it of $\delta v = 1$ cm ⁻¹ ; of	of $\lambda_R = 1$	$\mu = 0.00$	01 cm		
Orig.	δν	λ_R	Remarks ^b	Ref.	Orig.a	٥×	λ_R	Remarks	Ref o
rot.	60	160.7		Bo3,4	f	3221	3.105		Mag ²
	134	74.6		An		3224	3.102		CRi
	236	42.4		An		3225	3.101		BO
	140	71.4		Se		3228	3.10		HaC
	144	69.4		Hi ¹		3230	3.096		CC
	172	58.1		Bo3,4		3231	3.095		An
a	175	57.1		Mag ² , Hi ²		3233	3.093		Hu
-	200	50.0		Mag³, CBL		3246	3.081		Sp1
	340	29.4		De		3260	3.067		Sp^2
	440	22.7		Hi ¹		3270	3.058		Ro1
	464	21.6		An		3278	3.051		Sp1
а	500	20.0		Mag ^{2,3} , Hi ²		3290	3.04		Po
	510	19.6		Bo4		3324	3.008		$DuKo^{1,2}$
	550	18.2		CRi		3360	2.976		Ro1,2
	600	16.7		Mag1,2		3390	2.950		Ro1
	700	14.3		CRi, Mag ³		3394	2.946		Ki
a	740	13.5		Mag ²		3400	2,941		DeKg, Ry2
-	754	13.3		An		3405	2,937		Sp1,2
	780	12.8		Bo4		3406	2.936		Sp ¹
	1627	6.157		Hi ¹		3410	2.93	$(H_2O)_2$	Ro4
	1645	6.08		Bo^4		3419	2.925	, - ,-	DuKo1, Ro3
	1650	6.06		CRi		3420	2.924		HuC
	1656	6.04		CBL		3423	2.921		Ros
f	1659	6.03		Mag², Hi²		3428	2.917		An
	1665	6.00		An		3431	2.915		MeP
	1705	5.86_		KU		3433	2.913		Ro ^{6,7}
	2130	4.69		Mag ³	f	3435	2.911	Max	Bo ^{3,4} , HaC,
	2135	4.68		Mag^2					Mag_2
а	2150	4.65		Bo4, Hi2		3436	2.910		CRi, An
	2170	4.61		Hi ¹ , CBL		3437	2,909		N^2
		?) 4.25(?	')	GaV		3440	2.907		Ry¹, CBL
	3084	3.24	7	Ro ⁵		3443	2.904		Hu
	3100	3.226	1	Ry^2		3444	2.904		GhK
	3180	3.145	ļ	Ry^1		3445	2.903		Hi ¹
	3190	3.135	1	CBL		3448	2.900		Bh¹
	3195	3.130	H ₂ O	Bh¹,N²,Ki		3450	2.898		CC, CPR
	3199	3.126		GaV		3453	2,896		GaV
	3200	3.125	1	Bo3,4		3456	2.894		N_1
	3205	3.12	l	Ro4		3469	2.883		Bo
	3206	3.119	1	N^1		3474	2.878		KU
	3208	3.117	i	Ro3		3513	2.846		DKUo1,2
	3214	3.111	1	An		3554	2.814		Sp¹
	3217	3.108	Max	Ro8,7		3560	2.809		CC, Sp ²
	3219	3.106	I	Hi ¹		3569	2.802		Sp^{i}

Table 153—(Continued)

Orig.	δν	λ_R	Remarks	Ref.	Orig.	ε δν	λ_R	Remarks	Ref.
	3578	2.795	1	N^1		6042d	1.655d		ВО
	3582	2.792	Max	Ro3,4,6,7	c	6747	1.482	x	Hu
	3589	2.786	1	BO	c	7246	1.380	x	Hu
	3600	2.778		Ry2, An		7729	1.294		HuC
	3605	2.774		An		7757	1.289	x	Hu
	3609	2.771		GaV	c	8200	1.220	œ.	Hu
	3610	2.770	$(H_2O)_3$	Bh¹		8243	1.213		HuC
	3624	2.759		HaC	c	8660	1.155	x	Hu
	3625	2.758	1	CRi		8703	1.149		HuC
	3628	2.756	i	Ro ⁵	С	9175	1.090	x	Hu
	3630	2.755-	1	Bo3,4, Ry1		9223	1.084		HuC
	3650	2.740		CBL	С	9569	1.045		Hu
	3990	2.506		Bo4		10039	0.996		Hu
	4000	2.500		CBL		10151	0.985	oc	HuC
a	4023	2.486		Mag ^{2,3}	c	10635	0.940		Hu
С	5090	1.965		Mag ²		10944	0.910		Hu
	5100	1.961		Mag^2		11264	0.888	x	HuC
	5502	1.818		GaV					

^a Origin of the line: a = associated molecules (Mag^2) ; c = combination tones (Mag^2, Hu) ; f = fundamental frequency of the molecule (Mag^2) ; rot = from rotation of the molecule (Mag^2) .

^b Remarks: Square bracket indicates extent of band; Max = position of maximum, but DuKo, W. Gerlach ⁵⁴⁴ and Hu found in the band centered near 3400 cm⁻¹ only two maxima, which the first two place near 3324 and 3513 cm⁻¹, and Hu near 3233 and 3443. H₂O, (H₂O)₂, and (H₂O)₃ are the molecules to which I. R. Rao ⁵⁴⁵ assigns the associated values of δν. In that paper he replies to Su, who disagrees with him and has endeavored to interpret this band in terms of H₂O and (H₂O)₂ only; x indicates that Mag² reports that he did not find the line.

° References:

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 Bo1
BO
 CBL
                                                             Cross, P. C., Burnham, J., and Leighton, P. A., J. Am. Chem. Soc., 59, 1134-1147 (1937).
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 CC
CPR
CRi
 De
DeKa
   DuKo1
   GaV
 GhK
   HaC
   Hil
                                                           (1937).

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Meyer, E. H. L., and Port, I., Idem, 31, 509-510 (1930).

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                                                                  (1937).
   Hu
   HuC
   Ki
   KU
   Mag1
   Me1
   MeP
   NI
   Po
   Ro1
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⁶⁴⁴ Gerlach, W., Naturwissenschaften, 18, 68 (L) (1930).

⁵⁴⁵ Rao, I. R., Proc. Roy. Soc. (London) (A), 145, 489-508 (1934).

Table 153—(Continued)

Ry¹ Ramaswamy, C., Indian J. Phys, 5, 193-206 (1930); Ry², Nature, 127, 558 (1931). Segrè, E., Att. Accad. Lincei (6), 13, 929-931 (1931). Sp² Specchia, O., Nuovo Cim. (N. S.) 7, 388-391 (1930); Sp², Idem, 9, 133-137 (1932).

d These values are printed as 6642, 4.67, which are not self-consistent. It seems likely that there are two typographical errors; a 6 for a zero, and a 4 for a 1.

Table 154.—Abridged Raman Spectrum of Water

In this table the values that in the compiler's opinion may refer to the same band are connected by braces; in the case of the two prominent bands, centered near $\delta_{\nu}=1650~{\rm cm^{-1}}$ and 3400 cm⁻¹, only the extreme values and certain others of special interest are given, and the references in the "observer" columns apply to the band and not especially to the individual value on the line with them, except that the two "no's" on the line with $\delta_{\nu}=3610$ indicate that the maximum placed by several near 3610 was sought but not found by those observers. In the "observer" columns, r= recorded, no = sought but not found.

	•	0				()}		
Orig.b	δ×	λ_R	Bo3.4	CRi	Hu	Observera - HuC	Mag2-3	Misc
rot.	60	160.7	r					
	140)	71.4)						Sc
	144	69.4						Hi^{i}
	172)	58.1)	r					
a	175 🕻	57.1					r	
	200	50.0					r	$\mathrm{Hi^2}$
	340	29.4						De
	440	22.7						Hi ¹
	500)	20.0)					r	Hi^2
a	510 ≻	19.6	r					
	550	ر 18.2		r				
	550)	18.2)		r				
	600∫	16.75					r	
	700)	14.3		r			r	
a	740 }	13.5 }					r	
	780)	12.8)	r					
	1627	6.15						Hi¹
	1645)							
f	1659 }	6.03	r	r			r	KU, Hi ²
	1705							
	2130]	4.69]					r	
	2135	4.68					r	
a	2150 [4.65	r					Hi ²
	2170	4.61) 4.25 (?)						Hi ¹
	2355(?)	4.25 (7)						GaV
	3084	3.24)						
	3195	3.13						
f	3221	3.10	r	r	r	r	r	Many
	3400 ≻	2.94						
	3582	2.79						
	3610	2.77			no		no	
	3630	2.75)						
	3990 }	2.50 }	r					
a	4023	2.49 \					r	
C	5090 į	1.96)					r	
	5100 }	1.96				1	r	0.17
	5502	1.81						GaV

Table	154-	(Continued)

					- C	bserver•.		
Orig b	$\delta \nu$	λ_R	Bo2,4	CRi	Hu	HuC	Mag2.8	Misc
	6042°	1.66°						BO
C	6747	1.48			r		no	
c	7246	1.38			r		no	
	77291	1.291				r		
	7757	1.29			r		no	
C	8200)	1.22)			r		n)	
	8243 🕻	1.21				r		
c	86 60	1.12			r		no	
	870 3	1.15				r		
C	9175)	1.09)			r		no	
	9223	1.08∫				r		
C	9569	1.04			r			
	10039	1.00			r			
	10151	0.98				Г	no	
c	10635	0.94			r			
	10944	0.91			r			
	11264	0.89				r	no	

Observer: Misc = miscellaneous observers. For significance of the symbols designating the observers, see references in Table 153.
Origin of the line: See Table 153, note a.
See note d in Table 153.

Electron and β -ray Luminescence of Water.

High-speed electrons, such as β -rays, excite in water the same kind of luminescence as do y-rays (p. 304). Its intensity is not reduced by the common quenchers of fluorescence, nor by heating; it is partially polarized, the electric vector being parallel to the path of the electrons; its angular distribution is unsymmetrical, being much more intense in the direction of motion of the electron than in the reverse direction.546 I. Frank and I. Tamm 547 seek to explain this asymmetry in terms of electrons moving with velocities exceeding that of light in the medium (here water). Čerenkov 546, p. 105-108 reported the following relative values of the intensity (I) of the light emitted at an angle ϕ with reference to the direction of motion of the exciting electrons:

φ	0°	15°	30°	37.5°	45°	60°	75°	90°
Ί	63	68	73	37.5° 53	31	12.5	6.0	3.4

Mechanical Luminescence of Water.

In 1934 H. Frenzel and H. Schultes 548 announced that redistilled water luminesces under the action of ultrasonic vibrations, but that degassed water does not. L. A. Chambers, 549 using a frequency of 8.9 kc/sec observed such luminescence by 14 pure substances (including water) and some solutions, but none by 22 other pure substances. He has reported that the intensity of the luminescence varies "inversely with the temperature" and directly as $\mu\eta$, μ being the dipole moment and η the coefficient of

⁵⁴⁶ Cerenkov, P. A. (Tscherenkov), Compt. rend. Acad. Sci. URSS, 14, 101-105, 105-108 (1937). 547 Frank, I., and Tamm, I., Idem., 14, 109-114 (1937).

⁵⁴⁸ Frenzel, H., and Schultes, H., Z. physik. Chem. (B), 27, 421-424 (1934).

⁵⁴⁰ Chambers, L. A., J. Chem'l Phys., 5, 290-292 (1937) → Phys. Rev. (2), 49, 881 (A) (1937).

viscosity of the substance. There is no visible luminescence if $10^{18}\mu\eta < 1.94$ cgse. The light "originates in cavitated areas or at the surface of the cavities." Similar observations have been made by V. L. Levšin and S. N. Rževkin.⁵⁵⁰ They regard the light as an effect of the electrical potential differences that arise when the cavity is formed. They state that the light first appears at the liquid boundary, usually the lower, increases gradually in intensity and extent, until the entire volume of water is luminous, lasts for a time, and then abruptly vanishes.

40. Preparation of Dust-free Water

It is difficult to obtain a liquid free of suspended particles—exceedingly difficult in the case of water, much less difficult in the case of more mobile liquids. Various methods have been used and described in some detail by the workers here noted.⁵⁵¹

By taking extreme precautions, Lallemand attained partial success with distillation. Spring did not succeed with distillation, obtained some success with filtration through animal black, and success with both electrical separation (cataphoresis) and gelatinous precipitation (envelopment). obtained his best results by precipitation with Zn(OH)2, but found filtering through unglazed porcelain (Pukall filter) to be fairly satisfactory. Martin used repeated distillation in vacuo and without ebullition, fractional distillation in the same manner, envelopment, and cataphoresis. In his first paper, he reported that the remanent luminescence of water "is constant in intensity irrespective of the method of purification employed." In his second paper he stated that he believes this to be the first conclusive evidence for the scattering of light by pure substances. In that paper he also stated that the use of quartz vessels led to no improvement. Garrard has said that if there is no ebullition—no bubbling or bumping—during the distillation in vacuo, neither the actual temperature at which it is done nor the difference in the temperatures of the boiler and receiver affects the efficiency of the process for the removal of motes; that it is impossible to get dust-free water if the receiver contains either a piece of copper, or of vulcanized rubber, or a few cm³ of mercury; that if the receiver containing dust-free water is shaken, the water "is invariably contaminated again with motes," and that such recontamination is not prevented by shaking, rinsing back, and redistilling even as many as 20 times, and the same is true if the receiver is of quartz.

Sweitzer did not obtain satisfactory results with either ultrafiltration or centrifuging at 30 000 r.p.m., but did with envelopment. The least time for clearing when aluminum hydroxide was used as enveloper was two

⁸⁸⁰ Levšin, V. L., and Rževkin, S. N. (Lewschin and Rschevkin), Compt. rend. Acad. Sci. URSS, 16, 399-404 (1937).

⁵⁶¹ Lallemand, A., Ann. de chim. et phys. (4), 22, 200-234 (1871); Spring, W., Rec. trav. chim. Pays-Bas, 18, 153-168 (1899); Biltz, W., Nachr. Ges. Wiss. Gottingen (Math. Phys.), 1904, 300-310 (1904); Marain, W. H., Trans. Roy. Soc. Canada III (3), 7, 219-220 (1913), J. Phys'l Chem., 24, 478-492 (1920); Garrard, J. D., Trans. Roy. Soc. Canada III (3), 18, 126-127 (1924); Sweitzer, C. W., J. Phys'l Chem., 31, 1150-1191 (1927); Schade, H., and Lohfert, H., «Koll. Z., 51, 65-71 (1930); and Magat, M., Jour. de Phys. (7), 5, 347-356 (1934).

weeks. No impairment was observed to result from prolonged standing in Pyrex vessels. Schade and Lohfert used the methods employed with success by Spring and by Biltz, and also that employed by W. Gerlach⁵⁴⁴ i.e., repeated distillation from a copper vessel. They state that the last is not inferior to the others if proper precautions are taken. They found that quartz vessels are not suitable, that water standing in such vessels very soon gives evidence of containing particles in suspension, presumably on account of solution of the quartz. On the other hand, carefully cleaned vessels of hard Jena glass were entirely satisfactory. Magat used doubly distilled water filtered through collodion.

W. II. Martin and S. Lehrman 552 have stated that water distilled in lead glass scatters 50 per cent more light than that distilled in sodium glass, and that the results for sodium glass, for Pvrex, and for fused quartz are all alike.

See also the papers here noted. 553

DIFFRACTION OF X-RAYS BY WATER

When a slender pencil of x-rays is passed through water and impinges normally upon a photographic plate, the diffracted rays form upon the plate a principal dark ring, outside of which is a fairly uniform darkening which is rather sharply bounded along a ring concentric with the first. A more careful study has revealed 4 concentric rings at each of which the darkening of the plate passes through a maximum (see Table 155); but only the first is prominent. The intensities of these maxima, and to a less extent their positions, are affected by the absorption of the radiation by the water; W. Good 554 seems to have been the first to attempt to correct his data for water for this effect.

J. Thibaud and J. J. Trillat 555 have pointed out that, if the incident radiation contains both the general spectrum and the characteristic radiation, there will in general be two systems of rings, one arising from the characteristic radiation, and the other from the maximum of the general Their relative intensities will depend upon the thickness of the layer of water. For a copper target and 40 kv, both systems will show if the water is only 1 mm thick, but only the second if it is 8 mm.

Explanation of the rings is far from simple. 556 Early observers attempted to explain them either as arising from diffraction by neighboring, more or less polymerized, molecules 557 or as originating within the molecules. At that time, R. W. G. Wyckoff 558 inclined to the latter, but stated that the data "do not exclude the possibility of their arising from characteristic

⁵⁵² Martin, W. H., and Lehrman, S., J. Phys'l Chem., 26, 75-88 (1922).

⁵⁵⁸ Ananthakrishnan, R., Proc. Indian Acad. Sci., 2, 29-302 (1935); Magat, M., Ann. de Phys. (11), 6, 108-193 (1936); Mayer, J., and Pfaff, W., Z. anorg. allgem. Chem., 242, 305-314 (1935); and Malfitano, G., J. de Chim. Phys., 19, 32-33 (1921). The last gives some instructions for the preparation of collodion filters.

⁵⁵⁴ Good, W., Helv. Phys. Acta, 3, 205-248, 436 (1930).

⁵⁰⁵ Thibaud, J., and Trillat, J. J., Jour. de Phys. (7), 1, 249-260 (1930).

⁵⁸⁶ Amaldi, E., Physik. Z., 32, 914-919 (1931).

associations of molecules." C. V. Raman and K. R. Ramanathan 559 have developed a theory accounting for certain of the characteristics of the diffraction of x-rays by liquids on the basis of the fluctuations in density arising from thermal agitation. H. H. Meyer 560 was of the opinion that

Table 155.—Periodicities in the Diffraction of X-rays by Water

 $d = \lambda / \left(2 \sin \frac{\phi}{2}\right) = \text{equivalent grating space, } \phi = \text{angular deviation}$ corresponding to a maximum of the intensity of the diffracted rays, $\lambda =$ wave-length of the x-rays incident upon the water; $t \, {}^{\circ}C = \text{temperature}$.

When an author reports λ and d, but no ϕ , the values of ϕ , as computed from λ and d, are enclosed in parentheses.

I. A. Prins ⁵⁶⁴ has reported that inside the main ring (d = 3A) the darkening of the plate remains nearly constant until the place corresponding to d = 17A is reached, beyond which it decreases rapidly to a low limit.

More recently, a maximum between d = 4A and d = 5A has been observed, but it is not very pronounced. 565, 566

			Unit of A	and of $d =$	$= 1A = 10^{-1}$	$8 \text{ cm} = 10^{-}$	4 μ			
λ			,			a	!			Ref.a
0.712	13.44°	24°			3.04	1.71				ICT
1.54	29	46			3.07	1.97				ICT
1.54	27.3				3.25					So
1.539	(27.9)	(43.0)	(71.3°)		3.193	2.10	1.32		3	\mathbf{M}
1.539	(28.5)	(42.8)	(70.1)		3.130	2.11	1.34		20	\mathbf{M}
0.7090	(13.0)	(19.2)	(30.4)	(46.9°)	3.135	2.13	1.35	0.89	20	M
1.539	(28.8)	(42.8)	(66.7)		3.095	2.11	1.40		40	\mathbf{M}
					3.27	2.11				\mathbf{St}^{1}
					3.24	2.11	1.13		21	$\mathbf{St^2}$
1.54	30.5	41			2.93^{b}	2.20^{b}				G

- a References:
 - Good, W.554 Good, W. God. W. God. R. W. G., Int. Crit. Tables, 1, 338 353 (351) (1926). Values are based upon observations of Keesom, W. H., and DeSmedt, J., Proc. Akad. Wet. Amsterdam, 25, 118-124 (1922) [$\lambda = 0.712$]; 26, 112-115 (1923) [$\lambda = 1.54$]. Meyer, H. H., Ann. d. Physik (5), 5, 701-734 (1930). Sogani, C. M., Indian J. Phys., 1, 357-392 (1927). Stewart, G. W., Phys. Rev. (2), 35, 1426 (A) (1930); St², Stewart, G. W. God. Physical Conference of the physical Conference of the

After correcting for absorption. The other values in the table are not so corrected.

the main ring (d = 3A), see Table 155) arose from radiations scattered by adjacent molecules, and the next (d = 2.1) from those scattered by the

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Box Katzoff, S., J. Chem'l Phys., 2, 841-851 (1934).

⁸⁶⁶ Warren, B. E., J. Appl. Phys., 8, 645-654 (1937).

⁵⁵⁷ Keesom, W. H., and DeSmedt, J., Jour. de Phys. (6), 4, 144-151 (1923); 5, 126-128 (1924). 588 Wyckoff, R. W. G., Amer. J. Sci. (5), 5, 455-464 (1923). 559 Raman, C. V., and Ramanathan, K. R., Proc. Indian Ass. Cultiv. Sci., 8, 127-162 (1923).

⁵⁶⁰ Meyer, H. H., Ann. d. Physik (5), 5, 701-734 (1930).

⁵⁶¹ Stewart, G. W., Phys. Rev. (2), 37, 9-16 (1931).

Bes Debye, P., and Menke, H., Physik. Z., 31, 797-798 (1930).

⁵⁰⁸ Stewart, G. W., Phys. Rev. (2), 35, 726-732 (1930).

constituents of large complex groups of molecules; while G. W. Stewart 561 thought both of them arose from a single kind of molecular group. He advocated abandonment of the idea of an association into groups of a few molecules each, and its replacement by that of what he termed the cybotactic condition. In that condition, groups of hundreds or thousands of molecules have temporary existence, with ill-defined boundaries, and have a certain internal regularity. This accords with the conclusion of P. Debye and H. Menke, 562 that liquid mercury has a kind of quasi-crystalline structure. Evidence for the existence of the cybotactic condition in liquids has been summarized by G. W. Stewart. 563

Table 156.—Diffraction of X-rays by Water: Intensity and Effect of Temperature 560

I is the intensity relative to that (taken as 100) corresponding to d_1 at the same temperature, d = equivalent grating space.

	Un	it of $d=1$	$A = 10^{-8}$ cm.	Temp. =	1°C	
1	d_1a	I_1	d_2^a	I_2	da	I_3
3	3.193	100	2.10	21	1.32	5
20	3.130	100	2.11	18	1.34	5
40	3.095	100	2.11	14	1.40	5
Uncertainty	± 0.003		± 0.01		± 0.03	

°G. W. Stewart ⁵⁰¹ has reported that, as t increases, d_1 decreases at the rate of 0.0014A per 1 °C, d_2 increases, and at higher temperatures (above 40 °C) the maximum corresponding to d_2 vanishes. His observations extended from 2 °C to 98 °C. S. Katzoff ⁵⁰⁵ has reported that changing the temperature from 3 °C to 90 °C does essentially nothing to the diffraction pattern beyond reducing the prominence of its features. B. E. Warren ⁵⁰⁶ has stated that the distance between adjacent O's increases from 2.9A at 1.5 °C to 3.0A at 83° C. This corresponds to an increase of about 1 in 10 in the specific volume, whereas the actual change (Table 94) is only 3.1 in 100.

42. Absorption and Scattering of X-rays and of γ-rays by Water

Until quite recently, it was thought that in their passage through matter such high-frequency electromagnetic radiation as x-rays and γ-rays disturb the massive nuclei of the atoms but little, their direct effects being restricted to an interaction with the extra-nuclear electrons. But now it is known that they excite the nucleus, which subsequently emits certain characteristic radiations. L. H. Gray and G. T. P. Tarrant 507 have reported that this characteristic radiation from water consists of two components; for one, the coefficient of absorption in lead is $\mu_{Pb} = 0.85$ cm⁻¹; for the other, $\mu_{\rm Pb} = 1.96$ cm⁻¹.

The gross amount of energy expended in exciting the nuclei is, however, small as compared with that involved in the interactions between the radiation and the extra-nuclear electrons. Consequently, only the latter will be considered in the rest of this section, which accords with what has been the usual treatment of the subject.

The interaction between the radiation and an extra-nuclear electron may

⁵⁰⁷ Gray, L. H., and Tarrant, G. T. P., Proc. Roy. Soc. (London) (A), 136, 662-691 (1932).

Table 157.—Absorption of X-rays and of γ -rays by Water

The absorption by water of x-rays generated by 100 kilovolts is equivalent to that by lead that is only 0.004 as thick.⁵⁷¹ In the region $\lambda = 0.1$ to 0.5A, the relation between λ^3 and μ is not linear.⁵⁷² The apparent absorption (μ_a) of water for the radiation from Ra-C, perhaps affected by the presence of secondary radiation arising from the cosmic radiation, varies with the thickness (x) of the water as follows ⁵⁷³:

x	2	5	10	25 cm
1000 μ_a	17	25	28	38 cm ⁻¹

For a study of the variation in the quality and the intensity of x-rays as they pass through water, and of the way these vary with the size of the beam, see F. Vierheller.⁵⁷⁴

Except as the contrary is indicated, the following data have been taken from a compilation by J. A. Gray,⁵⁷⁵ which is based upon the work of J. Chadwick,⁵⁷⁶ J. Chadwick and A. S. Russell,⁵⁷⁷ C. W. Hewlett,⁵⁷⁸ A. R. Olson, E. Dershem, and H. H. Storch,⁵⁷⁹ F. K. Richtmyer,⁵⁸⁰ E. G. Taylor,⁵⁸¹ and K. A. Wingårdh,⁵⁸²

 $I = I_0 e^{-\mu x}$; $\mu_m = \mu M/\rho N_0$; $\rho = \text{density}$; M = formula-weight (H₂O) = 18.015; $N_0 = \text{number of molecules per g-mole} = 6.061 \times 10^{23}$; for the significance of other symbols, see text.

	Unit of $\lambda = 1A = 10$	$^{-8}$ cm; of $\mu = 1$ cm	n^{-1} ; of $\mu_m = 10^{-2J}$ cm	n² per molecule	
λ	μ	μm	λ	μ	μm
Cosmic ^a	0.00020	0.0006	0.340	0.290	0.861
Cosmica	0.00075	0.0022	0.360	0.309	0.918
Cosmic ^a	0.00157	0.0047	0.380	0.330	0.980
Cosmic a	0.00518	0.0154	0.400	0.352	1.05
Cosmica,	0.0183	0.054	0.420	0.376	1.12
0.00476	0.0437	0.130	0.440	0.400	1.19
Ra_{10}^c	0.0472	0.140	0.500	0.500	1.48
Rasd	0.0558	0.166	0.550	0.600	1.78
0.059_{1}^{d}	0.133	0.395	0.586	0.686	2.04
0.100	0.167	0.496	0.631	0.812	2.41
0.110	0.171	0.508	0.709	1.08	3.21
0.120	0.175	0.520	0.783	1.38	4.10
0.130	0.178	0.529	0.881	1.95	5.79
0.140	0.180	0.536	0.929	2.18	6.47
0.150	0.183	0.545	0.977	2.52	7.48
0.160	0.187	0.555	1.539	9.001	26.7
0.170	0.190	0.564			
0.180	0.194	0.576	Stumpen, H,	Physik Z., 50, 21	15-227 (1928).
0.190	0.198	0.588	0.158	0.186	0.55
0.200	0.201	0.597	0.211	0.204	0.60
0.220	0.212	0.630	0.264	0.238	0.70_{7}
0.240	0.223	0.662	0.317	0.280	0.832
0.260	0.234	0.695	0.370	0.335	0.99
0.280	0.246	0.731	0.423	0.384	1.141
0.300	0.259	0.769	0.475	0.482	1.43:
0.320	0.273	0.811	0.56	0.649	1.92,

^a The reported apparent coefficients of absorption (μ_a) of the ultrapenetrating "cosmic" radiations vary from 0.0002 to 0.0052.⁵⁸⁸

⁶1 Found by J. H. Sawyer ⁵⁸⁴ for the shower-producing cosmic rays.

^b γ-rays from Th-C", filtered through 6.8 cm of Ph. ⁵⁸⁵

Table 157—(Continued)

" γ -rays from Ra-B and Ra-C, filtered through 10 mm of Pb; value given is μ_a , the apparent coefficient.

^d Like the preceding, except that the filter is 3 mm of Pb. ^d₁ Reported by W. V. Mayneord and J. E. Roberts. ⁵⁸⁶

• Using a reflection method and nearly homogeneous x-rays of effective $\lambda = 0.14$ A, H. Fricke, O. Glasser, and K. Rothstein ⁸⁸⁷ found $\mu = 0.025$ 6.

Reported by H. Steps, 588 who used the Cu K_n-radiation.

result in (1) a transfer to the electron of the entire energy of the impinging quantum, thus destroying the radiation and removing the electron from its energy level; or (2) a transfer of a portion of the quantum energy to the electron, thus removing the electron and scattering the remaining energy of the quantum as a quantum of reduced frequency (Compton effect); or (3) an elastic impact in which the electron is not removed, but the direction of the path of the quantum is changed. All of these reduce the intensity of the transmitted beam.

If a unifrequent beam of parallel rays of such high-frequency radiation of intensity I_0 impinges normally upon a slab of material (the absorber) of thickness x, the intensity (I) of the transmitted beam, at a point far beyond the absorber and in the prolongation of the incident beam, is given by the formula $I = I_0 e^{-\mu x}$, where μ is called the coefficient of absorption. The coefficient μ is made up of three parts, each related to one or two of the three results already enumerated. One (τ) , called the coefficient of fluorescent or of photoelectric absorption, arises from the complete transfer of energy mentioned in result (1); another (σ_a) , called the coefficient of true absorption due to scattering, arises from the partial transfer mentioned in result (2); and the third (σ_s) , called the coefficient of true scattering, arises from the scattering or deviation of radiation mentioned in results (2) and (3). Thus $\mu = \tau + \sigma_a + \sigma_s$, which is often written $\mu = \tau + \sigma_s$ denoting $\sigma_a + \sigma_s$.

The vacancies left by the removal of electrons mentioned in results (1) and (2) are quickly filled, and that is accompanied by an emission of radia-

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and (2) are quickly filled, and that is accompanied by an emission of radia-

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Table 158.—Angular Distribution of the Radiation (x and γ) Scattered by Water

The various theoretical formulas for the distribution of the scattered radiation are all of the form $I_s = IAF^2(1 + \cos^2\theta)$, where I is the intensity of the incident radiation, A is the universal constant $e^4/2m_0^2c^4=3.96\times$ 10^{-28} cgs electrostatic units, F is a function of θ and of the structure of the atoms involved, and θ is the angle between the direction of propagation of the incident beam and that of the scattered beam of intensity I_s . For water, the values in the first section of this table, from a compilation by J. A. Gray, 589 were obtained by W. Friedrich and M. Bender, 590 the radiation being the platinum $K\alpha$ doublet, $\lambda = 0.19A$; they find for μ the very high value 0.236 cm⁻¹.

θ	(1+cos²θ)	$\left[egin{matrix} F_{ heta} \ F_{ heta 0} \end{matrix} ight]^2$	$\binom{I_{4,\theta}}{I_{4,90}}$	θ	(1+cos²θ)	$\begin{bmatrix} F_{\theta} \\ F_{90} \end{bmatrix}$	$\binom{I_{s,\theta}}{I_{s,0}}$
10°	1.97	1.50	2.95	90°	1.00	1.00	1.00
20	1.88	0.90	1.67	100	1.03	1.02	1.05
30	1.75	0.94	1.65	110	1.12	0.98	1.11
40	1.59	0.92	1.50	120	1.25	0.96	1.21
50	1.41	0.96	1.35	130	1.41	0.97	1.38
60	1.25	0.98	1.23	140	1.59	0.98	1.55
70	1.12	1.04	1.11	150	1.75	1.01	1.78
80	1.03	1.04	1.07	160	1.88	1.02	1.94
X-rays: λ=	0.31A. B	ackhurst, I	. Phil. Mag.	(7), 321–35	1 (1934).		
θ	30	40	50	60		70	80°
$I_{s,\theta}/I_{s,90}$	1.453	2.044	1.729	1.429	1.2	20	1.099
θ	100	110	120	130	14	40	150°
$I_{s,\theta}/I_{s,90}$	1.002	1.033	1.107	1.225	1.3	56	1.484

Table 159.—Coefficients of Scattering of X-rays and of γ-rays by Water

Adapted from a compilation by J. A. Gray.⁵⁹¹ It is not always possible to determine with certainty whether the coefficient found is that of the true scattering (σ_8) or that of the total scattering (σ) .

τ	Init of $\lambda = 1A = 10^{-8}$	em; of σ and $\sigma_0 = 1$	cm ⁻¹
λa	σε	σ	Source
Rac	0.0383		Neukirchen
0.161		0.185	Statz
0.240		0.206	Statz
0.285		0.170	Statz
0.32	0.198		Mertz
0.43	0.206		Mertz
0.501		0.201	Statz
0.54	0.210		Mertz
0.66	0.216		Mertz
0.79	0.228		Mertz

^e Effective wave-length of filtered x-rays, as defined by the equation: $(\mu/\rho)_{A1} = 14\lambda_{A}^{2.02}$, $\rho = \text{density}$; unit of $\mu/\rho = 1$ cm²/g, of $\lambda = 1$ A = 10^{-8} cm.

Sources:
 Mertz, P., Phys. Rev. (2), 28, 891-897 (1926); Neukirchen, J., Z. Phys., 6, 106-117 (1921);
 Statz, W., Idem, 11, 304-325 (1922).
 Gamma rays from Ra-B and Ra-C, filtered through 2.6 cm. of lead.

tion characteristic of the atom. In the long run, that emission is uniformly distributed in all directions and so might logically be called scattered radiation, but it is not so called.

The scattered radiation mentioned in (2) and (3), which is the only radiation technically described as scattered, is most intense in a direction that coincides, or nearly coincides, with the direction of propagation of the incident beam.

The intensity of the transmitted beam at points near the absorber is abnormally great on account of the presence of scattered radiations. At distances that are great as compared with the transverse dimensions of the absorber, the intensity of the scattered radiations decreases approximately as the inverse square of the distance from the absorber, while that of the transmitted beam, which by hypothesis is composed of parallel rays, is independent of that distance. Values of μ computed from measurements made very near the absorber will be called "apparent" coefficients of absorption and will be denoted by μ_a .

When γ -rays pass through water, the scattered rays showing the Compton effect have a wave-length about three times as great as that of the incident rays. Additional information on the quality of the radiation scattered by water when traversed by x-rays and by γ -rays is given by W. J. Rees and L. H. Clark 569; and H. Schindler 570 has studied the secondary radiation excited in water by the "cosmic" radiation.

43. Absorption and Transmission of Radiation by Water

(For x-rays, γ-rays, and cosmic radiation, see Section 42; for corpuscular radiation, Section 26; for scattering, Sections 39 and 47.)

The fraction of the incident radiation transmitted by a given layer of water depends upon the amount laterally scattered by the water, as well as upon the amount truly absorbed, i.e., converted into another form of energy. But the distinction has seldom been observed in reporting experimental data, the entire reduction in intensity being generally described as absorption. In some cases there is no necessity for maintaining the distinction, the scattering being experimentally negligible; but in other cases such is not the case.

Information regarding the scattering of radiation will be found in Sections 39 and 47, on luminescence and on the color of water, respectively;

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Chao, C. Y., Proc. Nat. Acad. Sci., 16, 431-433 (1930).
Mayneord, W. V., and Roberts, J. E., Nature, 136, 793 (L) 1935).
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Gray, J. A., Int. Crit. Tables, 6, 8-22 (17) (1929).
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that for absorption and transmission, with or without scattering, will be given here.

Pure Water. 592

The transparency of water is practically limited to the range $\lambda = 0.17$ to $1.0 \,\mu$. Within that range the transmissivity is great except near the limits, but outside it the absorption is very great until in one direction

Table 160.—Monochromatic Absorptivity of Water

(For x-rays and γ -rays see Table 157.)

The As, Dr, Kr, Ow, RA, and RL data have been taken directly from a compilation by J. Becquerel and J. Rossignol, who attribute the RL data to D. A. Goldhammer. The As values beyond $\lambda = 0.75 \,\mu$ correspond to the successive maxima and minima as reported by him; values in parentheses do not appear in the I.C.T.

A. Esau and G. Bäz ^{596a} have studied the absorption of water in the range $\lambda = 2.8$ to 10 cm, presenting the data graphically.

 $I = I_0 e^{-kx}$ where x is the length of path, in water, that corresponds to a reduction in the intensity of a beam of parallel rays from I_0 to I.

f.a→ ear→	Kr 1901	Кте 1901	Ts 1928	Ho 1933	DH 1934	Hab' 1935
¥λ			10			
98						
2.9			242°			
5.4			111			
6	58.4	68.8				93.0
6.2	0011	0010	86			,,,,
7.8			48			
0						30.0
1.6			20			
3	14.0	16.6				
3.5		2010	16 ^d			
8.0			9.4			
9.0			8.3			
0.0	7.9	9.0	7.3		8.0	2.95
3.0			5.1			
6.6			3.6			
8.4			6.0^{d}			
0	5.2	6.1				1.45
0.4			3.3			
4.6			4.6d			
8.8			4.1d			
0	4.8	5.7	3.6^{d}	6.4	3.3	0.86
0	2.9	3.4				0.47

502 For general reviews and discussions, see Schaefer, C., and Matossi, F., "Das Ultrarote Spektrum," 1930, Lecomte, J., Trans. Faraday Soc., 25, 864-876 (1929); Fowle, F. E., Smithsonian Misc. Collect., 68, No. 8 (Publ. 2484) (1917); Dawson, L. H., and Hulburt, E. O., Opt. Soc. Amer., 24, 175-177 (1934).

⁸⁹⁸ Leifson, S. W., Astroph. J., 63, 73-89 (1926).

⁸⁹⁴ Schaeffer, E. J., Paulus, M. G., and Jones, H. C., Physik. Z., 15, 447-453 (1914).

box Becquerel, J., and Rossignol, J., Int. Crit. Tables, 5, 268-271 (1929).

⁵⁰⁶ Goldhammer, D. A., "Dispersion und Absorption des Lichtes," Leipzig, 1913.

⁵⁰⁶a Esau, A., and Bäz, G., Physik. Z., 38, 774-775 (1937).

Table	160-	(Continued)
Table	100	COMMINUMENT

Ref. ^a → Year→ 10 ^a λ	Kr 1901	Kr. 1901	Ts 1928	Ho 1933	DH 1934	Hab' 1935
240 250	2.7	3.2	1	4.8	1.35	0.29 0.25
260 280	2.2	2.5		3.7 2.5	0.92 0.77	0.21 0.12
300	1.3	1.5		2.1	0.64	0.09
320 340				1.6 0.9	0.43 0.28	
360 380				0.9 0.9	0.19 0.13	
400				0.5	0.08	

For range $\lambda = 310$ to 800 m μ see Table 161.

Ref.•→ Year→ λ	As 1895	Co 1922	Dr 1924	Pl 1924	$\begin{array}{c} \operatorname{Ref}, \stackrel{a}{\longrightarrow} \\ \operatorname{Year} \rightarrow \\ \lambda \end{array}$	As 1895	Co 1922	Dr 1924	Pl 1924
0.745 0.845				0.044	1.45 1.475	(36.0)		20.1	
0.850 0.900•			0.069 0.0161		1.50 1.56	38.4		26.4 15.0	
0.950			0.0311		1.60	(21.0)		9.0	
0.970 0.980	0.446	0.448	0.480	0.142	1.677 1.708	(13.6) 11.4		5.2	
0.995 1.05	0.416		0.472 0.368		1.75 1.85	16.0		7.5 12.7	
1.085			0.333		1.90			31.5	
1.095 1.13 1.17	0.188 0.29		0.60			125		86	
1.20 1.21		1.22	1.12 1.30	1.28/	1.97 2.00 2.08	111 42	103	104 70 35.6	
1.243	1.22			1.20	2.10			31.6	
1.25 1.281	(1.21) (1.17)		1.24		2.147 2.15	27.8		24.7	
1.30 1.35	(1.20) (1.61)		1.48 2.14		2.237 2.30	(32)		19.6 25.9	
1.40 1.44	(23.4)	2.94	3.05		2.35 2.40	(61)		33.0 40.3	

$\operatorname{Ref}_{\lambda}^{a\rightarrow} \xrightarrow[\lambda]{} \operatorname{Ref}_{\lambda}^{a\rightarrow}$	As 1895	RL 1909	$\begin{array}{c} \operatorname{Ref}_{\cdot} \overset{\sigma}{\to} \\ \operatorname{Year} {\to} \\ \lambda \end{array}$	As 1895	RL 1909
2.6	(190)	530	6.09	2530	
2.8	•	2240	6.2	2060	2000
3.0		7330	6.5	1040	1030
3.02	2730		6.73	870	1000
3.2	(2590)	(6640)	6.765	880	
3.4		1440	6.92	820	
3.6		490	6.955	830	
3.93	204		7.0	(820)	810°
4.5	(447)	450	7.11	820	010
4.70	545		7.275	845	
5.27	308		7.41	790	
5.42	342		7.44	810	
5.47	335		7.49	800	
5.8	928	910	7.545	810	
6.0	2120	2140	7.65	765	

Table 160—(Continued)

Ref.e⊣ Year→ λ		As 1895	RL 1905		Ref. a→ Year→ λ	As 1895	k	RL 1909	
7.70 7.83 7.88 7.94 8.0	ŕ	785 765 775 690 (766)	755	;	8.38 8.43 8.49 9.0 10.0	695 755 725		700 703	
8.065 8.13 8.16 8.22 8.28		785 765 785 715 765			11.0 12 13 15			1200 2590 2890 3570 2990))
Ref. ^a → Year→ λ		RA 1898	RH 1910		RW 1911	Ow 1912		CE 1936	
24 52 61		>46 >46 >46						1160	
63 80			990)				1020	
83 100 108 117 152					460	423		710 460 360 320	
314						242		320	
			Unit of	λ=1 cm	; of $k = 1 \text{ cm}^{-1}$				
Ref.•→ Ec Year→ 1913	Te 1923	Za 9	Se Kn 33 1937	Rü 1918	Ref.a→ Ec Year→ 1913	Te Za 1923 1927	Se 1933	Kn 1937	Ra 1918
0.42 0.84 1.1 1.5 1.75 19.4	38.3 22.3 16.5 15.3	~			13.6 .14.0 14.48 15.29 18.41		0.38 0.40	0.47 0.45 0.31	

Year→ λ	1913	1923	1927	1933	1937	1918	Year→ 1913	1923	1927	1933	1937	1918
0.42 0.84 1.1 1.5 1.75	19.4	38.3 22.3 16.5 15.3	— к				13.6 .14.0 14.48 15.29 18.41		*	0.38 0.40	0.47 0.45 0.31	
1.8 2.7 3.7		16.1 10.5					19. 20.44	•	0.170	0.47	0.26	
4.80 5.34	5.84				3.53 2.92		23. 57.38 63.76		0.170			.0261 ^h .0228
5.7 6.48	2.38				2.11		67.98 74.80 84.68				0.	.0156 .0172 .0050
8.05 8.16					1.53 1.32		98.72 105.92				0.	.0057 .0099
8.80 9.55 10.10 10.87	0.86				1.17 0.99 0.90 0.76		128.81 144.12 144.50 159.66				0.	.0028 .0015 .0033 .0014
11.12					0.74		183.80 221.17				0.	00018
12.6 13.41				0.52	0.56		225.51 242.44				0.	00060

a References:

- CCS:

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 Cartwright, C. H., and Errera, I., Proc. Roy. Soc. (London) (A), 154, 138-157 (1936) ← Acta Physicochim. URSS, 3, 649-684 (1935) → Cartwright, C. H., Nature, 135, 872 (L) (1935); Idem, 136, 181 (L) (1935).

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 Haas, E., Biochem. Z., 282, 224-229 (1935) (Temp. = 24 °C).

 Hodgman, C. D., J. Opt. Soc. Amer., 23, 426-429 (1933).
- Co DH
- Dr Ec Ha

Table 160—(Continued)

Knerr, II. W., Phys. Rev. (2), 52, 1054-1067 (1937) \rightarrow Idem, 51, 1007 (A) (1937). Kreusler, H., Ann. d. Physik (4), 6, 412-423 (1901). Values as published in Int. Crit. Tables; see Kre. Kr as corrected by the compiler. Kn Kr

Kre

That Kreusler's published coefficients were seriously in error (by a factor of the order of 10) was pointed out by E. O. Hulburt, and Becquerel and Rossignol multiplied them by 10, getting the Kr values. By working backward from Kreusler's coefficients, through his recorded absorptions, the present compiler found that he had taken the length of the absorbing column as about 16.8 cm. For certain substances, he used a trough 16.85 cm long, but he states that he used for water a special quartz trough that was 20 mm long. Assuming that his reported absorptions are correct and that his trough was 20 cm long, we obtain the values given under Kre.

RA RH

RL

Owen, D., Electrician (London), 68, 504-507 (1912).
Plyler, E. K., J. Opt. Soc. Amer., 9, 545-555 (1924).
Rubens, H., and Aschkinass, E., Ann. d. Physik (Wied.), 65, 241-256 (1898).
Rubens, H., and Hollnagel, H., Verh. physik. Ges., 12, 83-98 (1910).
Rubens, H., and Ladenburg, E., Idem, 11, 16-27 (1909). The I.C.T. gives D. A.
Goldhammer ["Dispersion und Absorption des Lichtes," Leipzig, 1913] as the source Gothammer (Dispersion and Absorption des Lichtes, Delpzig, 1743) as and of these values.

Rückert, E., Ann. d. Physik (4), 55, 151-176 (1918).

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Seeherger, M., Ann. d. Physik (5), 16, 77-99 (1933).

Tear, J. D., Phys. Rev. (2), 21, 611-622 (1923).

Tsukamoto, K., Rev. d'Optique, 7, 89-108 (1928).

Zakrzewski, K., Bull. Int. Acad. Polon. Sci. Let. (A), 1927, 489-503 (1927).

Rü

RW Se

"The Ha values refer to 24 °C; absorption varies rapidly with the temperature, see Table 162. CO2 content is of slight effect.

S. W. Leifson 500 has stated that a thin film of water condensed on a window will absorb all radiation having $\lambda < 0.179 \,\mu$; see also H. Ley and B. Arends. See

° In the Ts paper the value for $\lambda = 0.1829 \,\mu$ is given as k = 4.72, which corresponds to only 1/10 of the reported transmission; the value (2.42) here given accords with that transmission.

⁴ For these values, Ts used commercial distilled water; for the others, redistilled, conductivity = $6(10^{-8})$ ohm⁻¹ cm⁻¹. The value (1.16) he gives for $\lambda = 0.1935 \,\mu$ is obviously wrong; his recorded absorption leads to k = 0.155, essentially 0.16.

• J. Kaplan 500 has published a curve showing the following maxima and minima:

0.88s 0.89_{7} 0.90 0.91 0.91_{5} 0.91_{8} 0.93 0.93 0.94 # λ 0.891000k15 9 15 9 15 14 25

Pl's values for $\lambda = 0.98$ and 1.21 are for water at 0 °C.

⁹ Given in the Becquerel and Rossignol compilation ⁸⁰⁵ as k = 890, but it was obtained from the RL data, and they lead to k = 808.

* From $\lambda = 50$ cm to $\lambda = 60$ cm there is much absorption, but G. Mie ** has found that there is no anomalous dispersion if the water is pure, and that traces of glass (?) in solution give rise to bands of the type reported by R. Weichmann. 601

 λ becomes of the order of a meter, and in the other it becomes of the order of 0.00005 μ , there being, however, a great gap $(0.17 \, \mu > \lambda > 0.00015 \, \mu)$ in which no data are available. A thin film of water condensed on a window absorbs all radiation for which $\lambda < 0.18 \,\mu^{593}$; a layer 1 cm thick absorbs 38 per cent of the radiation of $\lambda = 0.995 \,\mu$ ($k = 0.472 \,\mathrm{cm}^{-1}$), and 95 per cent of that of $\lambda = 1.4 \,\mu$ ($k = 3.05 \,\mathrm{cm}^{-1}$), and a layer only $10 \,\mu$ thick

⁸⁰⁷ Hulburt, E. O., J. Opt. Soc. Amer., 17, 15-22 (1928).

⁵⁰⁸ Ley, H., and Arends, B., Z. physik. Chem. (B), 4, 234-238 (1929).

⁵⁰⁰ Kaplan, J., J. Opt. Soc. Amer., 14, 251-256 (1927).

eoo Mie, G., Physik Z., 27, 792-795 (1926).

eon Weichmann, R., Ann. d. Physik (4), 66, 501-545 (1921) → Physik. Z., 22, 535-544 (1921).

absorbs 9.0 per cent of that of $\lambda = 2.8 \,\mu$ ($k = 2240 \,\mathrm{cm}^{-1}$). From $\lambda = 0.4$ to 0.52 μ the transparency is great.

It has been found that, whereas aqueous solutions of salts that do not form hydrates absorb nearly the same as does a layer of pure water of the same thickness as the water in the sample of solution under examination, those of salts that form hydrates absorb less than do such thicknesses of pure water.⁵⁹⁴

Table 161.—Monochromatic Absorptivity of Water in the Range $\lambda = 310$ to 800 m μ .

The As, Au, and Ma data have been taken directly from the compilation by J. Becquerel and J. Rossignol ⁵⁹⁵; the Ew data come from the same source, but each value has been multiplied by 2.303, so as to reduce the values to the basis used in this table. The value in parentheses does not appear in the I.C.T.

 $I = I_0 e^{-kx}$ where x is the length of the path, in water, that is needed to reduce the intensity of a beam of parallel rays from I_0 to I.

		U	nit of $\lambda =$	$1 \text{ m}\mu = 10$	-7 cm; of k	= 1 cm ⁻¹ .	Room	temp.		
Ref.a→ Year→ λ	Ew 1895	As 1895	Au 1904	Aue 1904	Pi 1918 ———— 10 ⁴	Ma 1922	Sa 1931	LS 1932	Ho 1933	DH 1934
310 313 320 325 300							840 690 580 512 461		1600	430
340 360 370 380							38? 281 200 148		900 900 900	280 190 130
400							72		500	80
415 420 430	81 74 53						46 41 30			
436 440	37					12	23			
450 460	28 25	20			34		18 15 15			
470 480 490	28 30 32	20	2				15 15 15			
494 500 510	35 37	20 22			30		16 17	200		
520 522	41	18	2	20	30		19			
530 539	44	8	3		22		21			
540 546	48	9	11			34	24			40
550	53	36	26			0.1	27	150		
557 558				56	36			,		

Table 161—(Continued)

Ref.a→ Year→	Ew 1895	As 1895	Au 1904	Aus 1904	Pi 1918 10	Ma 1922	Sa 1931	LS 1932	Ho 1933	DH 1934
λ 560	62	30	40		10	ж ———	30			
570	76	20	43				38			
578						64				
579		0.5	=0	78	56					
580	97	26	50							
589					96					140
590	161	78	89				85			
600	246	160	165	400	165		125			
602				188						
607				224						
610	272	190	220				160			
616				264	200					
618	285	212	240		206		170			
620 630	299	224	250				178 181			
	277	224	230		205		101			
636 640	315	235	275		2 2 5		200			
643	313	233	213	309			200			
648				309	236					
650	340	250	305		200		210			
658	0.10	200	000	339			2.0			
660	373	280	325	339						
663	313	200	020		245					
670	421	300			210					340
680	485	340								
690	575	400								390
700	690	550						450		070
710	890	790								
720	1080	1150								
730	1310	1750								
740		2300								
750		2410						3000		
775		(2410)								
800		2040								

G. Hüfner and E. Albrecht 602 reported the following values for bands of width Δλ centered on the indicated \(\lambda \). Temperature was 17 to 18 °C.

λ	449	468	487	506	527	552	576	602	631	664	$m\mu$
Δλ	6	6	8	8	8	11	11	18	18	13	$m\mu$
105k	28	27	49	4.3	53	76	114	250	282	394	cm ⁻¹

References:

As

Au

Aue

Aschkinass, E., Ann. d. Physik (Wied.), 55, 401-431 (1895). v. Aufsess, O. F., Ann. d. Physik (4), 13, 678-711 (1904). Au as corrected by DH. Dawson, L. II., and Hulburt, E. O., I. Opt. Soc. Amer., 24, 175-177 (1934). Supersedes Hulburt, E. O., Idem, 17, 15-22 (1928). Ewan, T., Proc. Roy. Soc. (London) (4), 57, 117-161 (1895). His published values of ε are defined by the relation $I = I_0(10)^{-4\pi}$, cf. Ewan, T., Phül. Mag. (5), 33, 317-342 (1892). The compiler has multiplied each of them by 2.303 so as to reduce them to the basis of this table. The values given in Int. Crit. Tables are incorrect, being ε , not k. Ew

them to the basis of this table. The values given in Int. Crit. Tables are incorrect, heing e, not k.
Hodgman, C. D., J. Opt. Soc. Amer., 23, 426-429 (1933).
Lange, B., and Schusterius, C., Z. physik. Chem. (A), 159, 303-305 (1932); 160, 468 (1932).
Martin, W. H., J. Phys'l Chem., 26, 471-476 (1922).
Martin, W. H., J. Trans. Wisconsin Acad. Sci., Arts, Let., 19, 562-593 (1918).
Sawyer, W. R., Contrib. Canadian Biol. (N. S.), 7, 75-89 (No. 8) (1931). Ho LS

⁶⁰² Hüfner, G., and Albrecht, E., Ann. d. Physik (Wied.), 42, 1-17 (1891).

Table 162.—Absorptivity of Water: Effects of Pressure and Temperature

J. R. Collins ⁰⁰⁸ has found that changing the pressure from 120 atm to 5000 atm produces no change in the absorption of water in the range $\lambda = 0.71$ to 1.05μ .

He has also found 604 that an increase in temperature shortens the wavelengths at which the absorption has maxima, and changes the coefficients of absorption corresponding to the maxima. E. Ganz 605 has confirmed Collins' observations on the band at $\lambda = 0.77 \,\mu$. He seems to state that the coefficient of mass absorption (k/ρ) for any fixed λ varies linearly with the temperature for each of the bands $\lambda = 0.77 \,\mu$ and $\lambda = 0.84 \,\mu$, ρ being the density of the water; but he may mean that k so varies.

Using $\lambda = 12.6$ cm, M. Seeberger ⁶⁰⁶ found that the absorption decreases rapidly as the temperature rises.

 $I = I_0 e^{-kx}$.

Unit of $\lambda_{max} = 1\mu$; of k_{max} and k = 1 cm⁻¹, temp. = t °C.

I. J. R. Collins. 604

•	- 00	0.4-	05.00
			95 0
λmax	k max	Δλmax	$\Delta k_{ m max}$
0.740	0.0380	-0.035	0.0100
0.845	0.0472		
0.970	0.606	-0.015	0.176
1.17	1.38	-0.04	0.10
1.43	28.7	-0.02	-1.1
1.94	108	-0.02	0
	0.740 0.845 0.970 1.17 1.43	0.740 0.0380 0.845 0.0472 0.970 0.606 1.17 1.38 1.43 28.7	λωαν kmax Δλωαν 0.740 0.0380 -0.035 0.845 0.0472 0.970 0.606 -0.015 1.17 1.38 -0.04 1.43 28.7 -0.02

II. M. Seeberger. $\lambda = 12.6$ cm.

III. E. Haas. 607 Illustrative. He concludes that this increase in k is not due to the increase in the dissociation of the water.

1000λ→	18	6 ———	190 k	
17	0.670	0.690	0.280	0.295
37	0.930		0.420	0.420

Table 163.—Total Transmissivity of Water

As the absorptivity of a substance varies with the frequency of the radiation, numbers expressing the total transmissivity are of significance only with respect to a specified source of radiation and to a specified thickness of the substance.

In the following, τ is the transmissivity for radiant energy, and τ_l is that for light. If I and I_0 are the energies of the transmitted and of the incident radiation, respectively, and if L and L_0 are the corresponding luminous

oon Collins, J. R., Phys. Rev. (2), 36, 305-311 (1930) → 35, 1433 (A) (1930).

⁶⁰⁴ Collins, J. R., Idem, 26, 771-779 (1925).

⁶⁰⁵ Ganz, E., Ann. d. Physik (5), 26, 331-348 (1936) ← Diss., München, 1936.

⁶⁰⁶ Seeberger, M., Ann. d. Physik (5), 16, 77-99 (1933).

⁶⁰⁷ Haas, E., Biochem. Z., 282, 224-229 (1935).

Table 163—(Continued)

intensities, then $\tau = 100 \ I/I_0$ and $\tau_l = 100 \ L/L_0$. The temperature of the radiating source is $t \, {}^{\circ}\text{C} = T \, {}^{\circ}\text{K}$; the thickness of the transmitting layer of water is x.

Unit of x=1 cm; of $\lambda=1\mu=10^{-4}$ cm; of τ and $\tau_I=1\%$

I. Ideal radiator: "Black-body."

	$-x = 0.05 \text{ (Br)}^{\circ} -$		
7		ı	•
2.9		720	15.2
4.0		730	15.3
5.3		760	15.7
9.0		810	18.1
9.9		820	18.7
10.3		870	20.6
11.4		933	22.8
13.3		940	23.0
13.1		960	23.6
13.8			
1 (IC	CT)a		10 (ICT)a
7	71	7	TI
		Q	98.96
50	99.92	31.5	99.16
69	99.93	54	99.26
	2.9 4.0 5.3 9.0 9.9 10.3 11.4 13.3 13.3 13.8 7 22.5	7 2.9 4.0 5.3 9.0 9.9 10.3 11.4 13.3 13.8 13.8 13.8 22.5 99.90 50 99.92	729 4.0 720 4.0 730 5.3 760 9.0 810 9.9 820 10.3 870 11.4 933 13.3 940 13.8 7 1 (ICT) ^a 7 7 22.5 99.90 50 99.92

II. Various sources.

<i>x</i> ·		0.1	0.2	0.5	1.0	2.0
Source	T			τ (FC)a		
Iron	1000	3.1	1.4	0.5	0.25	0.10
Carbon	2150	35	28	20	15	11
Tungsten	2970	66	59	51	43	36
Sun		85	81	76	71	65
(We)a			$x\rightarrow$	0.0019		0.0038
Source	Source Filter		λο	T -		
Welsbach mantle None			107			20.0
Mercury arc	None			55.5	5	33.0
Mercury arc 2 mm quartz			60.3		38.4	
Mercury arc Cardboard		310	62.7		39.8	

" References:

Br Brown, S. L., Phys. Rev. (2), 21, 103-106 (1923).
FC Forsythe, W. E., and Christison, F. L., J. Opt. Soc. Amer., 21, 150 (1931).
ICT Compilation by Walsh, J. W. T., and Buckley, H., Int. Crit. Tables, 5, 264-268 (1929), based upon Aschkinass, E., Ann. d. Physik (Wied.), 55, 401-431 (1895), v. Aufsess, O. F., Diss., Munich, 1903

Ann. d. Physik (4), 13, 678-711 (1904), and Ewan, T., Proc. Roy. Soc. (London) (A), 57, 117-161 (1894).

We Weniger, W., J. Opt. Soc. Amer., 7, 517-527 (1923).

Table 164.—Penetration of Solar Radiation into Water

Computed by W. Schmidt 608 on the basis of the coefficients of absorption as determined by Aschkinass, 609 and the distribution of energy in the solar spectrum. All values for the radiation refer to energy; none to luminosity.

^b Corresponds to a tungsten filament vacuum lamp burning at 9 lumens per watt.

[&]quot;Corresponds to a plain carbon arc.

^d Approximately noon sun-light.

^{*}This is the wave-length of the principal radiation.

⁶⁰⁸ Schmidt, W., Sitz. Akad. Wiss., Wien (2A), 117, 237-253 (1908).

⁶⁰⁰ Aschkinass, Ann. d. Physik (Wied.), 55, 401-431 (1895).

Table 164.—(Continued)

A= fraction of total incident radiation absorbed by a layer of water of thickness x, and lying within the indicated range in wave-length. For example: The amount of radiation in the range $\lambda=0.9$ to $1.2\,\mu$ that is absorbed from sunlight by a column of water 1 cm thick is 5.60 per cent of the total solar energy incident upon the surface of the column.

 τ = the rate at which the temperature of an exceedingly thin layer of water at the depth x would rise if it retained in itself all the solar energy that it absorbs. (The incidence is presumably normal.)

			Unit of	$\lambda = 1\mu = 0$.001 mm;	of $\tau = 1$ °C pe	er min.		
$\stackrel{x \to}{\lambda}$	0.01mm	0.1mm	1mm	1cm	10cm	000A 1m	10m	100m	
0.2 to 0.6	0.00	0.00	0.00	0.1	0.8	7.5	65 0	223.1	237.0
0.6 to 0.9	0.00	0.04	0.7	6.3	54.8	230.1	350.2	359.7	359.7
0.9 to 1.2	0.08	0.67	6.8	56.0	170.6	178.8	178.8	178.8	178.8
1.2 to 1.5	0.54	4.78	23.3	69.5	86.6	86.6	86.6	86.6	86.6
1.5 to 1.8	1.82	16.28	53.0	80.0	80.0	80.0	80.0	80.0	80.0
1.8 to 2.1	2.00	14.05	25.0	25.0	25.0	25.0	25.0	25.0	25.0
2.1 to 2.4	0.84	6.42	24.2	25.3	25.3	25.3	25.3	25.3	25.3
2.4 to 2.7	0.94	5.24	7.2	7.2	7.2	7.2	7.2	7.2	7.2
2.7 to 3.0	0.19	0.40	0.4	0.4	0.4	0.4	0.4	0.4	0.4
Sum	6.41	47.88	140.6	269.8	450.7	640.9	818.5	986.1	1000.0
7	6.68a	3.69	0.71	0.071	0.0071	0.0008	0.00008	0.000003	

Radiation Filters Containing a Layer of Water.

A cell 0.5 mm thick containing distilled water and provided with quartz or fluorite windows transmits essentially no radiation beyond the approximate range $\lambda=0.17~\mu$ to 1.5 μ , and throughout that range the transmission is high except near the limits. Increasing the thickness restricts the range but slightly.

In his compilation, K. S. Gibson⁶¹⁰ gives the spectral ranges that may be isolated by the use of such water filters, either alone or in combination with other filters. He cites the following users of water filters: W. W. Coblentz,⁶¹¹ cell 10 mm thick with thin quartz windows; T. Lyman,⁶¹² cell 0.5 mm thick with fluorite windows, and cell 20 mm thick with quartz windows; H. Kreusler,⁶¹³ cell 20 mm thick with quartz windows.

Natural Waters.

" Surface layer.

The transmission of light by many coastal and inland waters is subject to wide fluctuations caused by variations in the turbidity and in the plankton. The amount of plankton varies with the season and the weather; and the turbidity with the amount of detritus, sand, and soil, whether brought in by streams or surface drainage, or stirred up from the bottom. Measurements of the transmission and of the effective absorptivity under such conditions are of no general value, but are of significance with reference to the

⁶¹⁰ Gibson, K. S., Int. Crit. Tables, 5, 271-274 (1929).

⁶¹¹ Coblentz, W. W., Sci. Papers Bur. Stand., 17, 725-750 (S438) (1922).

e19 Lyman, T., "Spectroscopy of the extreme ultra-violet," 1914.

⁶¹³ Kreusler, H., Ann. d. Physik (4), 6, 412-423 (1901).

actual plankton growth at the place and time considered; many such measurements have been made for the purpose of obtaining data for correlating plankton growth with the illumination existing at various depths. 614

A few illustrative sets of measurements on waters of this kind will be found in Table 166.

O. F. v. Aufsess ⁰¹⁵ has published curves showing the spectral absorption of the waters from several lakes. He was of the opinion that every departure of the color from that of pure water is due to the presence of foreign bodies, and that variations in the turbidity of a given lake change its color but little.

Table 165.—Monochromatic Absorptivity of Sea-water

The following data refer to samples taken from the open sea far from land, and at the depths indicated; the effect of scattering is probably small. E. O. Hulburt ⁶¹⁶ studied samples from the Pacific, the Gulf Stream, and the Caribbean Sea, and could detect no difference in their absorptivities. Tsukamoto concluded that bromides are the cause of the great absorptivity for $\lambda < 220$ m μ . The earliest study of the ultraviolet absorption by seawater seems to have been by J. L. Soret. 617

 $I = I_0 e^{-kx}$; I is the intensity of a parallel beam of radiation at a distance x along the beam beyond the point where the intensity is I_0 , both points being in the water.

Unit of $k=1$ cm ⁻¹ ; of depth =	= 1 meter; of $\lambda = 1$ m $\mu = 0.001 \mu = 10$ A	Room temp.
---	--	------------

	K. Tsukamoto ^a				E. O. Hulburta			
Depth→	Surface	3m - 1000k -	15m	Depth→	Surface	Tapb 1000k _	Dist ^b	
212.3	1079		1247	254	154	104	79	
213.6	646			266	131	74	48	
216.1	160		361	280	90	46	35	
217.5	145		206	303	39	16	12	
217.9		267		313	21	7	5	
221.0		58		366	3.0	2	2	
221.8	120		185	436	0.23		0.12	
226.4	74		137	546	0.35		0.341	
227.6		42		578	0.7		0.640	
231.7		44		612	2.3		2.3	
233.6		70	118					

a References:

Hulbert, E. O. 507
Tsukamoto, K., Compt. rend., 184, 221-223 (1927).

Tap = water supply of Washington, D. C.; Dist = doubly distilled water. Values for Dist and $\lambda = 436, 546, 578$ were taken from W. H. Martin, and $\lambda = 612$ from O. F. v. Aufsess. 615

⁶¹⁴ Regnard, P. M. L., J. Chem. Soc. (London), 60, 2 (1891) ← Mém. Soc. Biol. (Paris), 42, 288 + (1890) → Compt. rend. Soc. Biol. (9), 11, 289 (1890); Shelford, V. E., and Gail, F. W., Publ. Puget Sound Biol. Sta., 3, 141-176 (1922) (Bibliog. of 29 entries); Poole, H. H., Sci. Proc. Roy. Dublin Soc. (N. S.), 18, 99-115 (1925); Poole, H. H., and Atkins, W. R. G., J. Marine Biol. Assoc. United Kingdom (N. S.), 14, 177-198 (1926); 15, 455-483 (1928).

⁶¹⁵ v. Aufsess, O. F., Ann. d. Physik (4), 13, 678-711 (1904) ← Diss., München. 1903.

⁶¹⁶ Hulburt, E. O., J. Opt. Soc. Amer., 13, 553-556 (1926).

⁶¹⁷ Soret, J. L., Jour. de Phys. (1), 8, 145-158 (1879) \leftarrow Arch. des sci. phys. et nat., 61, 322-359 (1878); 63, 89-112 (1878) \rightarrow Compt. rend., 86, 708-711 (1878).

ons Martin, W. H., J. Phys'l Chem., 26, 471-476 (1922).

Table 166.—Effective Absorptivity of Some Coastal and Inland Waters

Illustrations of the variability of such data when obtained under natural conditions.

 $I = I_0 e^{-kx}$. Layers are specified by the depths of their bounding planes below the surface of the water.

Unit of $\lambda = 1$ m μ ; of k = 1 cm⁻¹

I. Various Waters. C. D. Hodgman.a

Waterb→ λ	Dist	Erie	Тар	Shaker 10	00kBr	Br'	Marsh	Snow
220	6.4	134	28	150	150	99	99	101
240	4.8	92	5.3	106	92	60	69	58
260	3.7	67	4.8	78	69	44	55	46
280	2.5	53	4.8	60	53	34	46	39
300	2.1	41	3.0	48	39	32	41	34
320	1.6	34	2.1	39	30	25	30	30
340	0.9	28	0.9	30	23	21.4	20.0	28
360	0.9	20.7	0.5	21.4	17.1	18.7	16.4	23
380	0.9	14.3	0.5	14.3	11.1	15.7	13.1	20.7
400	0.5	10.6	0.5	8.1	6.4	7.6	9.9	17.7

II. Gunflint Lake, Minn. H. A. Erikson.^a

$\lambda \rightarrow$	447	466	500	529	545	563	581	623
100k	1.046	0 740	0.501	0.357	0.357	0.322	0.309	0.407

III. Sea, total depth = 9 m; photographic. Martin Knudsen.

λ→ Layer	400	450	500	00k	600	650
1 to 3 m	0.58	0.30	0.16	0.19	0.30	0.38
3 to 5 m	0.40	0.27	0.21	0.20	0.28	0.38
5 to 7 m	0.64	0.39	0.27	0.38	0.49	0.60
1 to 8 m		0.300	0.184	0.181	0.275	

IV. San Juan Archipelago. Photronic cell with filters B, G, and R; B transmits from $\lambda=410$ to $\lambda=500$, max. at 465; G from $\lambda=500$ to $\lambda=590$, max. at 540; R uniformly from $\lambda=600$ to $\lambda=700$. C. L. Utterback and J. W. Boyle.^a

Filter→ Layer	B Stat	G 100k ion 1. k con	R	Filter→ Layer	B	G 100k — Station 4.	R
1 to 6 m	0.248 Stat	0.201 ion 2. k con	0.435	0 to 5 m 5 to 10	0.232 0.215	0.187 0.190	0.456 0.396
50 to 20 m	0.338	0.273 Station 3. 0.207	0.483	10 to 15 15 to 20 20 to 25	0.187 0.213 0.233	0.171 0.168 0.202	0.336 0.362 0.262
10 to 15 15 to 20	0.275 0.280	0.208 0.223	0.378 0.356	25 to 30 30 to 35	0.216 0.163	0.211 0.183	0.209
20 to 25 25 to 30 30 to 35	0.179 0.172 0.165	0.210 0.206 0.188	0.212 0.169	35 to 40 40 to 45 45 to 50	0.156 0.095	0.184 0.138 0.089	

Table 166—(Continued)

Alaskan coastal waters. Apparatus as for preceding. C. L. Utterback.a

Filter→ Layer	В	G 100k	R	Filter→ Layer	В	G 100k	R
	,	Station H	•	,	′	Station R	
0 to 10 m	0.165	0.126	0.407	0 to 5 m	0.322	0.231	0.427
10 to 15	0.068	0.120	0.414	5 to 10	0.579	0.231	0.457
15 to 20	0.087	0.124	0.287	10 to 15	0.391	0.351	0.575
20 to 25	0.248	0.116	0.104	15 to 20	0.091	0.410	0.271
25 to 30	0.400	0.113	0.051	20 to 25	0.061	0.234	0.095
30 to 35	0.322	0.118		25 to 40		0.110	
		Station SS				Station CS	
0 to 5 m	0.558	0.397	0.563	0 to 5 m	0.166	0.168	0.410
10 to 20	0.256	0.172	0.370	5 to 10	0.164	0.169	0.408
20 to 25	200	0.171		10 to 15	0.161	0.167	

a References:

Erikson, H. A., J. Opt. Soc. Amer., 23, 170-177 (1933). Hodgman, C. D., Idem, 23, 426-429 (1933). Knudsen, M., Cons. perm. intern. l'explor. mer, Publ. de Circons No. 76, 1922. Utterback, C. L., J. Opt. Soc. Amer., 23, 339-341 (1933). Utterbach, C. L., and Boyle, J. W., Idem, 23, 333-338 (1933).

Waters studied:

Dist = distilled water; Erie = Lake Erie; Tap = water supply of Cleveland, Ohio, drawn from Lake Erie; Shaker = Shaker Lake, Ohio; Br and Br' = two brooks; Marsh = water from an open marsh; Snow = melted snow.

Table 167.—Penetration of Daylight into Sea-water (Cf. Table 164.)

I. Absorptivity. Otto Krümmel.^a Based on observations of Regnard (citation not given) using a selenium cell; expressed in form $I = I_0 e^{-kx}$, where x is depth beneath surface. Unit of x = 1 m; of k = 1 m⁻¹.

II. Visual brightness; $B_h = looking horizontally$, $B_v = looking verti$ cally upward, B_h observed by W. Beebe and G. Hollister. a $B_{h'}$ and $B_{v'}$ computed b by E. O. Hulburt, a assuming the total illumination of the surface to be 10,000 candle/ $ft^2 = 10.764$ ca/cm.² Unit of B = 1 ca/cm²; of depth = 1 ft.

Depth	$10^3B_{\rm A}$	103Bh	108 Bv	Depth	10^3B_h	102Bh	$10^a B_v$
0			10764	300	0.17	0.17	17
50	11.4	11.3	1140	350	0.13	0.125	13
100	3.6	3.66	366	500	0.024	0.066	6
200	0.4	0.44	44	800	0.024	0.014	1
250	0.23	0.23	23				

Spectral distribution of visual brightness at a depth of 800 ft. in the sea, the brightness at the surface being i_0 ; brightness looking horizontally = b_h , looking vertically upward = b_v ; unit is arbitrary. Values of b, and b, as computed by E. O. Hulburt a for sea-water containing no suspended particles. Unit of $\lambda = 1 \text{ m}\mu = 0.001 \mu$.

Table 167—(Continued)									
λ	śo	104ba	102b.	λ	io	104ba	102b,		
400	2.10	0.00	0.00	500	4.27	7.40	7.20		
420	2.87	0.00	0.00	520	4.25	2.54	3.68		
440	3.55	0.158	0.115	540	4.19	0.09	0.27		
460	4.08	2.30	1.64	560	4.10	0.00	0.00		
480	4.24	6.54	5.12	580	3.97	0.00	0.00		
500	4.27	7.40	7.20	600	3.83	0.00	0.00		

IV. Spectral composition of light in the sea. Derived by W. R. G. Atkins ⁶¹⁹ from photographic data obtained by Grein in the Mediterranean Sea in 1913 and 1914. Designation of colors: R = red, OY = orange-yellow, G = green, BG = blue-green, B = blue, BV = blue-violet. $I \equiv I_c/I_t$, I_c and $I_t = \text{intensity}$ of the indicated color and total intensity, respectively, both at the same depth (d). Unit of d = 1 m.

$Color \rightarrow d$	R	Ο¥	G 10	001 BG	В	BV
1 5	96.7	165.7	165.7	165.7	198.9	207.3
	0.98	1.18	117.3	117.3	254.4	508.8
10 20	$0.34 \\ 0.018$	1.06 1.05	89.64 4.68	89.64 17.26	$\frac{282.2}{279.7}$	537.1 697.2
50	0.0025	0.069	4.53	5.04	486	504
75		0.054	4.73	14.2	193.6	787.5
100 200		0.0052	1.56 3.18	1.73 8.06	346.2 37.16	650.8 952
500			12.27	30.63	30.8	920.3
1000				74.6	37.31	881.1
1500						(1000)

- V. Yearly means from photoelectric measurements. H. H. Poole and W. R. G. Atkins. 620
 - (a) No regular seasonal changes in opacity.
 - (b) Level of maximum absorption changes during the day, presumably from migration of zoöplankton.
 - (c) Mean value of k in $I = I_0 e^{-kd}$ for the layers 0 to 20, 20 to 40, and 40 to 60 m, are 0.150, 0.120, and 0.111 m⁻¹, respectively.
 - (d) For most turbid, k = 0.228 m⁻¹; for least, k = 0.06 at depth of 25 m.
 - (e) The percentage of the incident light that reaches a depth of 20 m is .6.62, 40 m is 0.72, and 60 m is 0.085.
 - (f) Down to 25 m, the horizontal illumination is 0.54 of the vertical.
- (g) A Secchi disk is just visible when its illumination is 16 per cent of that of the surface of the water.

* References:

Beebe, W., and Hollister, G., Bull. N. Y. Zool. Soc., 33, 249-263 (1930). Hulburt, E. O., J. Opt. Soc. Amer., 22, 408-417 (1932). Krümmel, O., "Handb. d. Ozeanog.," Vol. 1, 1907.

^b Computed on the assumption that the true coefficients of absorption as determined by Hulburt are correct, that scattering is caused by thermal fluctuations in the concentration of the molecules, and that in the first 250 ft there is turbidity equivalent to one mote 0.1 mm² in sectional area in each cm³ of water. The last is introduced in order to make the computed values fit Beebe's observations.

44. Emissivity of Water

For radiation of wave-length greater than 1 μ , the absorptivity of water is very great (Section 43) and the reflectivity is small (Table 146); whence one may conclude that water will radiate very nearly as an ideal (black body) radiator, and that its emissivity will not be much less than unity. An experimental determination is difficult, and until recently there was only that of K. Siegl, 621 which for some reason leads to a value that is surely too low. In contrast to that, E. Schmidt 622 found that $E=0.985\pm0.001$ and does not vary with the temperature of the water, which was varied from 10 °C to 50 °C. Here E is the ratio of the radiation from water to that from an ideal radiator at the same temperature (Table 288). He found that a layer of water 0.1 mm thick radiates as strongly as does a thick layer.

45. Photoelectric Effects for Water

Two types of photoelectric effect are exhibited by water: one is photovoltaic, and the other has to do with the emission of electrons from an illuminated surface of water.

Photovoltaic Effect for Water.

R. Audubert 623 observed that if one of two metal plates immersed in an aqueous solution of an electrolyte is illuminated and the other not, the two plates acquire a difference in electrical potential. His investigation of this phenomenon forms the subject of a series of papers. 624 He concludes that water is essential to the phenomenon, and that it is photolyzed, *i.e.*, separated into O and OH, by the light.

Photoelectric Emission by Water.

Using a mercury arc shining through a fluorite window, W. Zimmermann 625 found that the illumination of a clean, fresh, water surface resulted in no emission of photo-electrons, but as the surface aged an emission analogous to that from solids appeared and grew. This he attributed to dust deposited from the air.

On the other hand, W. Obolensky, 626 using a spark between terminals of aluminum, observed an emission that varied with the filtration of the light, and that occurred only for wave-lengths shorter than about $\lambda = 200 \,\mathrm{m}_{\mu}$. (See Table 168.) Likewise, L. Couson and A. Molle 627 found an

⁶¹⁹ Atkins, W. R. G., J. du Cons. Int. Expl. Mcr., 7, 171-211 (1932).

⁶⁵⁰ Poole, H. H., and Atkins, W. R. G., J. Marine Biol. Assoc. United Kingdom (N. S.), 16, 297-324 (1929).

ea Siegl, K., Sitz. Akad. Wiss Wien (Abt. IIa), 116, 1203-1230 (1907).

⁶²² Schmidt, E., Forsch. Gebiete Ingenieurw., 5, 1-5 (1934).

eas Audubert, R., Compt. rend., 189, 800-802 (1929).

⁶²⁴ Audubert, R., Compt. rend., 189, 1265-1267 (1929); 193, 165-166 (1931); 194, 82-84 (1932); 196, 475-478, 1588-1590 (1933); Jour. de Phys. (7), 5, 486-496 (1934); Audubert, R., and Lebrun, G., Compt. rend., 198, 729-731 (1934).

⁶²⁵ Zimmermann, W., Ann. d. Physik (4), 80, 329-348 (1926).

emission when the exciting light lay in the range $\lambda = 200 \text{ m}\mu$ to $130 \text{ m}\mu$, and concluded that it was not due to casual contamination of the surface. They suggested that these electrons come from the oxygen in the water molecule.

More recently, P. Görlich, 628 using an evacuated monochromator with an optical train of fluorspar, has confirmed the observations of Obolensky and of Couson and Molle. He has shown that the limiting wave-length at which the emission vanishes as λ is increased lies between 203 and 204 m μ ; that it is independent of the actual value of the low conductivity of the water, which in his tests was varied from 3.3×10^{-6} to $3.2 \times 10^{-4} (\text{ohm cm})^{-1}$; and that it is the same for concentrated aqueous solutions of AgNO₃, NaCl, Na₂SO₄, or K₄Fe(CN)₆ as it is for pure water. He also studied the variation of the emission with the wave-length of the incident light (see Table 168).

In contrast to the preceding results stand those of H. Greinacher. 629 He used an electron counter, a flowing jet of water, and radiation from a quartz mercury-lamp, the filtration being 3 mm of fused quartz. Radiation so filtered is supposed to contain no wave-length shorter than 220 m μ , yet he reported a marked emission of electrons. The water contained air and lime.

Table 168.—Photoelectric Emission by Water

The ICT data were obtained with light from an Al-spark, filtered as indicated and containing no wave-length shorter than λ_{\min} ; the intensity of the emission by CuO when illuminated by the fluorite-filtered radiation corresponded to S = 40,000.

For the Görlich data, I is the intensity of the emission by water when illuminated by monochromatic light of wave-length λ and of a certain arbitrarily fixed intensity.

	Unit o	$f \lambda = 1 \text{ m} \mu = 10 \text{A} = 10$	⁻⁷ cm	
	ICTa		Görli	cha
Filter	\lambda min	S	λ	1
Glass	330	0	204.16	0
Calcite (CaCO ₃)	220	0	203.0	4.0
Quartz and air	177	11	198.9	17.5
Quartz (SiO ₂)	145	15	189.9	50.0
Fluorite (CaF2)	125	100	185.4	52.0
			176.3	21.4
			171.9	11.1

[&]quot; Sources:

ICT From the compilation by A. L. Hughes, Int. Crit. Tables, 6, 67-69 (1929); based on Obolensky, W., Ann. d. Physik (4), 39, 961-975 (1912).
 Görlich, P., Idem, (5), 13, 831-850 (1932).

^b H. Greinacher ^{exp} observed an emission when $\lambda > 220 \text{ m}\mu$ (see text). ^c Fluorite with not more than 3 mm of air at atmospheric pressure.

⁶²⁶ Obolensky, W., Ann. d. Physik (4), 39, 961-975 (1912).

⁶²⁷ Couson, L., and Molle, A., Arch. Sci. Phys. et Nat. (5), 10, 231-242 (1928).

⁶²⁸ Görlich, P., Ann. d. Physik (5), 13, 831-850 (1932).

⁶⁰⁰ Greinacher, H., Helv. Phys. Acta, 7, 514-519 (1934).

46. THE SPECTRUM OF WATER (For Raman spectrum, see Section 39.)

Under-water Sparks.

The spectrum of the light from an electric discharge (spark, arc, etc.) occurring in water consists of a continuous, and frequently intense, background extending far into the ultraviolet; of lines characteristic of the electrodes; in some cases, of the spectrum of water-vapor; usually of lines due to hydrogen, but those due to oxygen are absent except in the brush discharge, where they have been observed by H. Smith. B. Setna^{630a} has reported the presence of the following "water bands" in the spectra of under-water sparks: $\lambda = 7760$, 7933, 8226, 8475A. See also ⁶³¹.

A brief report on the emission of light by spark discharges in liquids has been made by J. A. Anderson,⁶³² in which other additional references are given. The subject has been studied primarily for the purpose of ascertaining how the spectrum of the electrode material is changed by the high pressure that exists in such discharges.

Absorption Spectrum.

The absorption spectrum of water at a given temperature is independent of the recent thermal history of the water, being the same for water from recently melted ice as for that from recently condensed steam.⁶³³

Table 169.—Absorption Spectrum of Water

The approximate position of other, generally less pronounced, bands may be determined from the data in Tables 160 and 161, and from the curves published by RL^a and Re.^a

 $\lambda_{\rm obs}$ and λ_m are, respectively, the observed value of the wave-length of greatest absorption and the mean of the better determinations of that wavelength; $\nu_{\rm obs}$ and ν_m are, respectively, the reciprocals of $\lambda_{\rm obs}$ and λ_m . Only the bands of pronounced absorption are represented in the λ_m column; and the values there given have been taken from ${\rm El}^{2,a}$

The width and intensity of a band are in some cases indicated by letters, b = broad, n = narrow, st = strong, wk = weak.

		τ	Init of $\lambda = 1\mu = 10^{-4}$	cm; of $\nu = 1$ c	m ⁻¹ .		
λ_m , (ν_m)	λ_{obs}	$\nu_{\rm obs}$	Ref.	λ_m , (ν_m)	λobe	Pobs	Ref.
0 55 (18200)	0.55 0.599 ^b 0.635 0.60	18200 16690 15750 16700	As MRB MRB RuLa	0.75 (13300)	0.745° 0.762 0.768 0.77	13420 13120 13020 12980	El', Pl Ga Ma As
0.63 (15900)	0.655 ^b 0.656 0.670 0.700 ^b	15270 15240 14920 14280	MRB MRB MRB MRB, RuLa	0.85 (11760)	0.842 0.845* 0.89 0.90	11880 11830 11200 11100 •	Ga El', Ma, Pl Ka Ka

680 Smith, H., Phil. Mag. (6), 27, 801-823 (1914).
680a Setna, B., Indian J. Phys., 6, 29-34 (1931).
681 Bloch, L. and E., Compt. rend., 174, 1456-1457 (1922); Finger, II., Verh. physik. Gcs.,
11, 369-376 (1909); Z. wiss. Photog., 7, 329-356, 369-392 (1909); Konen, H., Ann. d. Physik (4),
9, 742-780 (1902); Konen, H., and Finger, H., Z. Elektrech., 15, 165-169 (1909); Liveing, G. D.,
and Dewar, J., Phil. Mag. (5), 38, 235-240 (1894); Toriyama, Y., and Shinohara, U., Nature, 132,
240 (1933).

Table 169—(Continued)

λ_m , (ν_m)	λ_{obs}	₽obs	Ref.	λm, (νm)	λ_{obs}	vobs.	Ref.ª
7. meş (P me)	0.915	10930	Ka	2.97 b, st	2.97	3370	Pa
	0.913	10800	Ka	(3370)	3.03	3300	ES
	0.973	10280	La	(0010)	3.04	3290	Re
0.98	0.98	10200	Ma, Pl		3.06	3270	As, RL
(10200)	0.995	10050	Dr		3.08	3250	Er PIC
(10200)	1.00	10000	As		3.30	3030	PIC
1.18	1.20	8330	Ma		4.66	2146	Re
(8470)	1.21	8260	Dr	4.70 wk	4.70	2128	As, Ch, RL, ES
(0410)	1.215	9230	Pi	(2128)	4.72	2119	Pa
	1.25	8000	As	(/	5.56	1798	PIC
	1.44	6940	Pl		5.83	1715	PIC
	1.445	6920	ES		5.83 6.05	1653	Re
	1.45	6900	Do		6.06	1650	Pa
1.46	1.47	6800	Ma		6.08	1645	RL
(6850)	1.475	6780	Dr. St	6.1 n, st	6.10	1639	As, AF, Cb ES
(0000)	1.48	6760	Gr	(1639)	6.20	1613	ES
	1.50	6670	As, Cb	, ,	6.8	1470	Wi
	1.50 1.51 1.74 1.79	6620	Pa		7.1	1410	AF, RL
	1.74	5750	Ela		7.3	1370	AF
	1.79	5590	El³, Co		8.2	1220	AF
	1.93	5180	Es		8.6	1160	Wi
	1.94	5150	Ās		8.8	1140	AF RL
	1.95	5130	Ma		9.5	1050	RL
	1.954	5120	St		10.4	960	AF
	1.96	5100	Do, ES		12.5	800	AF AF, RL
	1.97	5080	Dr, Do		15.8	633	AF, RL
1.98n	1.98	5050	Gr		19.5^{d}	513	E13
(5050)	2.05	4880	Cb, Pa		20.	500	Ca AF
(4000)	2.79d	3580	El ³		20.2	495	AF
	2.90d	3450	El ³		60.	167	Ca
	2.94	3400	Er		83.	120	CaE
	2.95	3390	Cb, Ma, St		2000	5.0	Te
		2070	,,		7000	1.4	Te
					20000/	0.5	Te

Region 2 cm to 28 cm is still to be investigated. Between 28 cm and 300 cm there are probably no bands (Kn).

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⁸³² Anderson, J. A., Int. Crit. Tables, 5, 433 (1929).

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Table 169—(Continued)

b Sharp edge of band.

° Pl's observations refer to 0 °C.

^d These three values are used by El³ in his interpretation of the spectrum; the 2.79 and 2.90 μ represent 2 of the 3 Raman-spectrum components corresponding to the 3 μ band; the 19.5 μ band "is somewhat hypothetical" and interpreted as arising from the mutual vibrations of two (H2O)'s.

These Pa values varied with the thickness of the water. For greater values of λ , see Tables 177 and 178.

Table 170.—Analyses of the Absorption Spectrum of Water

Uncertainties in the values of the wave-lengths corresponding to the maxima of the several bands, and the complexity resulting from the overlapping of bands (see first section of this table), make the analysis of the spectrum difficult. Furthermore, there are differences of opinion as to the manner in which the vibrations of a molecule are affected by its neighbors. Some hold that the fundamental vibrations of the isolated molecule (those characteristic of the vapor) play no part in the case of the liquid, being completely suppressed by the action of neighboring molecules (e.g., Errera, 1937 a; Bosschieter and Errera, Compt. rend., 204, (1937 a)); some hold that the liquid is a mixture of polymers, each having fundamental vibrations of its own (Ellis, 1931 a; Cabannes and de Diols, 1934 a; Rao, 1934 a); and others hold that the fundamental vibrations of the isolated molecule remain predominant but, perhaps, restricted and with frequencies more or less modified, and that to these must be added additional fundamental vibrations arising from the bonding of molecules into a more or less definite, though transient, structure (e.g., Magat, 1936^a).

Until there has arisen some generally accepted opinion regarding these matters, the original articles should be consulted by those interested in the interpretation of the absorption spectrum of water. Some of the recent or more typical of those articles are listed in footnote a to this table.

As illustrative of the analyses that have been proposed, those by Ellis and by Magat are here given. They are the most detailed. Ellis first interpreted in terms of two fundamentals ($\lambda = 2.97$ and 6.1μ) all the values in column λ_m of Table 169, except $\lambda = 4.70 \,\mu$. But in order to obtain that and the two bands that he discovered at $\lambda = 1.79$ and 1.74μ , he replaced the $\lambda = 2.97 \,\mu$ by two others ($\lambda = 2.90$ and $2.79 \,\mu$) and included a fourth fundamental ($\lambda = 19.5 \mu$). The last is somewhat doubtful, and the two that replace the $\lambda = 2.97 \,\mu$ are derived from the Raman spectrum (see notes to Table 169). Magat first interpreted the spectrum in terms of three fundamentals ($\lambda = 2.9$, 3.1, and 6.0 μ), and later added 4 other fundamentals arising from the interaction of adjacent molecules. All five of these analyses are given below. In Magat's analyses n_1 , n_2 , n_3 have the same values in both cases, and are regarded as corresponding respectively to the σ , π , and δ (see Table 64) vibrations of the free molecule. In Ellis' analyses the frequencies have been so numbered as to correspond as closely as may be with the similarly numbered ones in Magat's.

Table 170—(Continued)

In all cases, the calculated frequency (cv) of a band is given by the relation $\nu = n_1\nu_1 + n_2\nu_2 + \ldots$, where $\nu = 1/\lambda, \nu_1 = 1/\lambda_1, \ldots, \lambda_1, \lambda_2, \ldots$ being the wave-lengths of the assumed fundamental vibrations, the n's being small integers, and c being the velocity of light. The values of the n's corresponding to each of the calculated λ 's are tabulated; the fundamental λ 's are those for which one n is unity and all the others are zero. $I = \inf_{n \to \infty} I$ red; R = Raman spectrum; "single" = only one of the n's occurs in the calculation of λ_{calc} ; "comb." = both n's occur. Except in the first of Ellis' analyses, all the n's listed occur in the calculation, each with its indicated value.

Unit of $\lambda = 1 \ \mu = 10^{-4} \ \text{cm}$; of $\nu = 1 \ \text{cm}^{-1}$ Ellis (1927)° Single Comb. Comb. 712 λobe 112 113 722 **n**3 0 0 0 6.1 1 6.1 2.97 1 2 0 0 3.05 2.97 1.98 0 3 1 1 2.03 1.96 2 0 4 1 2 1.52 1,45 1.48 1.46 5 2 1.22 1.17 1.18 23 3 6 2 2 3 0.98 0.98 0.97 1.015 7 0.85 0 0.8750.840.75 4 2 0.765 0.725 0.73 9 Õ 0 0 0.675 0.63 5 10 0.58 0 0 0.61 0 0.55 0.555 11 Ellis (1931)^a 12 Acaic λobs 111 116 19.5 0 0 0 1 19.5 6.1 0 0 0 6.1 1 0 0 4.71 1 4.65 2.90 0 0 1 0 2.90 0 0 2.79 2.79 1 0 1.79 0 1 1 1 1.785 1.74 1.745 -Ellis and Sorge (1934)^a λobs 721 752 77.3 Peale 6.20 0 0 0 6.21 1 1610 4.70 0 0 0 1 4.69 2130 0 0 2 0 3220 3.11 3.11 0 2.92 0 0 2.92 3430 1 2.80 1 0 0 0 2.80 3570 0 1 1 0 1.98 5040 1.93 1 0 1 0 1.93 5180 ō 0 0 2 1 1.80 5560 1.78 1 0 1 1.75 5700 Ō Ō 1.47 0 1,46 6860 1.42 2 0 0 0 1.39 7170 20 0 1 0 1.18 8470 1.18

1.14

8750

Table 170—(Continued)

, ·				— ма	agat (1	934) 4				
I	— Aobs —	R	n 1		n2		n 3	λ	eale	Veale
2.95 1.95 1.48	6. 3. 2.	028 105 911 965	0 0 1 1		0 1 0 0		1 0 0 1	3. 2. 1.	028 105 911 964 502	1659 3221 3435 5092 6660
1.20 0.98 0.84 0.77			1 3 3 3		1 0 0 1		1 0 1 0	0. 0.	.203 .970 .836 .736	8310 10310 11960 13590
				— Ма	agat (1	936) 4				
	R	111	n2	n 3	14	ns	116	227	Acale	Veale
19.6 14.9 6.02	167 66–44 20.0 13.5	0 0 0 0	0 0 0 0	0 0 0 0 1	0 0 0 1	1 0 0 0 0	0 0 1 0	0 1 0 0	167 60.2 19.6 14.3 6.02	60 166 510 700 1660
5.85 5.40 4.684 3.14 2.90	4.684	0 0 0 0	0 0 0 1	1 1 0 0	0 0 0 0	1 0 0 0	0 0 1 0	0 1 0 0	5.81 5.49 4.63 3.05 2.898	1720 1820 2160 3280 3450
2.486 1.95 1.789 1.47 1.18	2.485	1 1 1 1	0 0 0 1 1	0 1 1 0 1	0 0 0 0	0 0 0 0	1 0 1 0 0	0 0 0 0	2.53 1.957 1.779 1.486 1.192	3950 5110 5620 6730 8390
0.98		1 3 1 3	2 0 2	0 0 1	0 0 0	0 0 0	0 0 0	0 0 0	1.011 0.966 0.866	9890 10350 11550
0.84		3	2 0 3	î 0	0	0	Ŏ 0	0	0.833 0.763	12010 13110
0.75		3	1 3	0 1	0	0	0	0	0.73 4 0.677	13630 14770
0.66		3	1 3	1 2	0	0	0	0	0.654 0.609	15290 16430

a References:

Bosschieter, G., and Errera, J., Compt. rend., 204, 1719-1721 (1937); 205, 560-562 (1937); Jour. de Phys. (7), 8, 229-232 (1937). Cabannes, J., and de Riols, J., Compt. rend., 198, 30-32 (1934). Carrelli, A., Nuovo Cim. (N. S.), 14, 245-256 (1937). Cartwright, C. H., Nature, 136, 181 (L) (1935); Phys. Rev. (2), 49, 470-471, 421 (A)

(1936).

(1936).
Cartwright, C. H., and Errera, J., Acta Physicochim. URSS, 3, 649-684 (1935); Proc. Roy. Soc. (London) (A), 154, 138-157 (1936).
Ellis, J. W., Phil. Mag. (7), 3, 618-621 (1927); Phys. Rev. (2), 38, 693-698 (1931).
Ellis, J. W., and Sorge, B. W., J. Chem'l Phys., 2, 559-564 (1934).
Errera, J., J. de Chim. Phys., 34, 618-626 (1937).
Kinsey, E. L., and Ellis, J. W., Phys. Rev. (2), 49, 105 (L), 209 (A) (1936).
Magat, M., Jour. de Phys. (7), 5, 347-356 (1934); Ann. de Phys. (11), 6, 108-193 (Bibliog. of 148) (1936).

Piekara, A., Acta Phys. Polon., 6, 130-143 (1937). Rao, I. R., Proc. Roy. Soc. (London) (A), 145, 489-508 (1934). Williams, D., Phys. Rev. (2), 49, 869 (A) (1936).

Table 171.—Effect of Temperature and Pressure on the Absorption Spectrum of Water

J. R. Collins ⁶³⁷ found no change in the absorption spectrum of water when the pressure was increased from 120 atm to 5000 atm, the range covered being $\lambda = 0.71 \,\mu$ to $1.05 \,\mu$.

Collins (1937)^a found no marked change in the band at 1.79 μ when the temperature was increased from 4 to 137 °C. Ganz (1936)^a has reported that the intensity of the 0.77 μ band increases with the temperature, that the band near 0.84 μ is scarcely detectable at room temperature, but is well-defined and sharp at 87 °C, and (1937)^a that the band at 4.7 μ is still visible at 84 °C.

Of historical interest only are the early observations of W. J. Russell and W. Lapraik ⁶³⁸ who were unable to detect visually any change in the absorption spectrum when the water was heated from 20 °C to 60 °C.

 λ = wave-length at which the absorption is a maximum for the band considered.

	Unit of $\lambda =$	$1 \mu = 10^{-6} \text{ cm}$		
Collins (1925)" -		S	tansfeld (1925) "
λ — 95 °C	Decr. in λ	20 °C	λ — 97 °C	Decr.
0.740 0.845	0.035	1.475 1.954	1.468	0.007 0.009
0.970 1.17	0.015 0.04	1.701	1.710	0.007
1.43 1.94	0.02 0.02			
	95 °C 0.740 0.845 0.970 1.17 1.43	follins $(1925)^a$ Decr. $\ln \lambda$ 0.740 0.035 0.845 0.970 0.015 1.17 0.04 1.43 0.02	λ — Decr. in λ 20 °C 0.740 0.035 1.475 0.845 1.954 0.970 0.015 1.17 0.04 1.43 0.02	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$

[&]quot; References:

Collins, J. R., Phys. Rev. (2), 26, 771-779 (1925); 52, 88-90 (1937). Ganz, E., Ann. d. Physik (5), 26, 331-348 (1936) = Diss., München, 1936; Idem, 28, 445-457 (1937). Stansfeld, B., Z. Physik, 74, 460-465 (1932).

47. THE COLOR OF WATER AND OF THE SEA *

Pure Water.

Transmitted light.—It is generally stated that long columns of pure water appear blue by transmitted light. Lord Rayleigh 630 has written that W. Spring somewhere stated that in columns 4 or 5 meters long the color of pure water is a fine blue, only to be compared with the purest skyblue as seen from a great elevation; but that when the incident light was white he himself has never obtained "a blue answering to Spring's description."

- W. Spring 640 is positive that the blue color pertains to the water itself,
- *W. D. Bancroft 642 has published a review and summary of many of the earlier papers treating of this subject.
 - 687 Collins, J. R., Phys. Rev. (2), 36, 305-311 (1930) → Idem, 35, 1433 (A) (1930).
 - 628 Russell, W. J., and Lapraik, W., J. Chem. Soc. (London), 39, 168-173 (1881).
 - coo Lord Rayleigh, Proc. Roy. Inst. Grt. Brit., 19, 765-771 (1910) = Nature, 83, 48-50 (1910).
 - 640 Spring, W., Rec. trav. chim. Pays-Bas, 18, 1-8, 153-168 (1899).
 - 641 Spring, W., Ibid., 17, 359-367 (1898).

and that the presence of suspended matter modifies that color, introducing a greenish tint, and in some cases rendering the water colorless, as in some lakes. A very small amount of exceedingly small particles of hematite in suspension will suffice for the last.⁶⁴¹

From a study of the waters from a number of lakes, O. F. v. Aufsess ⁶¹⁵ concluded that every departure from the blue of pure water arises from the presence of foreign substances, and that the color of a given lake is only slightly affected by changes in the turbidity.

In contrast with the preceding we have: (a) the statement of J. W. Lovibond 643 that a 4-foot stratum of distilled water is equivalent in color to a combination of the Lovibond filters Yellow 1.0 and Blue 1.45, which color is an unsaturated green; and (b) the fact that the values of the absorptivities given in Table 161 indicate that the color of a stratum of water 20 meters or less in thickness is very definitely green-blue, quite similar in hue to that of the spectrum near $\lambda = 0.49 \, \mu.^{643a}$ It should be remembered that none of the water used in obtaining the data in Table 161 was really optically empty, even the best being only so dust-free as may be obtained by repeated distillations (cf. Section 40), and in some cases no correction was made for the effect of the windows.

It has been suggested that the blueness of water arises from the relatively few molecules of ice that are dissolved in the water. Such was the opinion of J. Duclaux.⁶¹⁴ He regarded pale green as the color of hydrol itself, and suggested that a study of the variation in the color of water with the temperature might yield data from which the relative proportions of the two polymers (ice and hydrol) could be computed. This is in line with Barnes' statement that the color of the St. Lawrence River changes as freezing becomes imminent (see Section 58).

Scattered light.—A. Turpain ⁶⁴⁵ has called attention to a series of papers published some 60 years ago by Alexandre Lallemand, and apparently forgotten. Lallemand ⁶⁴⁶ observed that most, probably all, liquids scatter light laterally, even when devoid of suspended particles. He called this fluorescence, and suggested that it arose in part from a kind of molecular reflection or diffusion, and in part from free vibrations of the molecules. That arising in the first way will be partially or completely polarized and of the same frequency as the incident light exciting it; that arising in the second will be unpolarized and of a different (he says longer) wave-length. He attributed sky light also to molecular scattering.

C. V. Raman ⁶⁴⁷ has shown that the intensity of the light molecularly scattered by a liquid, and the attendant coefficient of absorption, can be calculated on the "theory of fluctuations"—the theory that the number of

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Bancroft, W. D., J. Franklin Inst., 187, 249-271, 459-485 (1919).
Lovibond, J. W., "Light and Colour Theories," p. 21, 1915.
Gibson, K. S., and Keegan, H. J., J. Opt. Soc. Amer., 27, 58 (A) (1937).
Duclaux, J., Rev. gén. des Sciences, 23, 881-887 (1912).
Turpain, A., Compt. rend., 197, 1107-1109 (1933).
Lallemand, A., Ann. de chim. et phys. (4), 22, 200-234 (1871); (5), 8, 93-136 (1876).
Raman, C. V., Proc. Roy. Soc. (London) (A), 101, 64-80 (1922).
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molecules per unit of volume varies slightly from point to point, on account of the thermal agitation of the molecules. The intensity so calculated for dust-free (optically empty) water is 160 times that of dust-free air; an observed value was 175 times. Likewise the computed coefficients of absorption for $\lambda = 0.494$ and 0.522μ , where there is negligible selective absorption, are 0.000029 and 0.000022 cm⁻¹, respectively; while the experimental value found by Aufsess for each was 0.00002 cm⁻¹ (see Table 161).

He states that a sufficiently thick layer of pure water exhibits by molecular scattering a deep blue color more saturated than sky light and of comparable intensity. "The colour is primarily due to diffraction, the absorption only making it of a fuller hue."

The light scattered laterally by thin columns of water comes mainly from suspended particles, and is not blue, partaking largely of the color of the incident light as modified by the selective reflection of the particles.⁶⁴⁰ Spring held the erroneous opinion that there is no scattering by the water itself.

The deep blue color of Crater Lake, Oregon, probably arises from the scattering by the water itself, modified, more or less, by the color of the skv.648

For methods employed for obtaining optically empty water, see Section 40.

The Sea.

Although long columns of pure water are blue by transmission, and the light scattered by thick layers of water is blue, there has been much discussion regarding the blueness of the sea. Three effects may contribute to the color: (1) the reflection of the blue sky; (2) the proper color of the water, including the effect of molecular scattering; and (3) the scattering of small particles held in suspension.

Lord Rayleigh 639 attributed most of the color to the reflection of the sky, and thought that only a very unimportant fraction of it is to be accounted for by the scattering by small particles. Whether he intended the last to include molecular scattering is not entirely clear. To these conclusions, C. V. Raman 647 and J. Y. Buchanan 649 have taken exception, attributing the blueness to the water itself. Buchanan based his objection on his observation that when quiet water, as in the screw-well of the Challenger, is viewed vertically under such conditions as to exclude reflected sky, it appears to be of a beautiful dark-blue color. Truly, as Rayleigh remarked in reference to the water at Capri, this light, scattered or internally reflected by the water, came in large part from the sky. would enhance the blueness if the sky were blue, but it can scarcely be accepted as the sole explanation of the observations recorded by Buchanan. In reality, the discussion seems to have arisen, at least in part, from differ-

Pettit, E., Proc. Nat. Acad. Sci., 22, 139-146 (1936).
 Buchanan, J. Y., Nature, 84, 87-89 (1910).

ences in the interpretation of the vague term "color of the sea." Does it mean the color seen when the surface of the sea is viewed from above and at such a distance that the line of sight makes a large angle with the vertical, there being nothing to obstruct the reflection of the sky from the portion observed? If so, the intensity of the reflected light will be so great, as compared with that coming from the interior of the water, that the observed color will depend almost entirely upon that of the sky. This is what Rayleigh seems to have had in mind. But if the term "color of the sea" means the color of the light coming directly from its interior, unmixed with light reflected from the surface, then the color of the sky will play a very subordinate part. Under intermediate conditions, the relative importance of each of those sources of color will vary with the condition. Again, one may mean by "color of the sea" the color of the light transmitted by a very long column of sea-water when the incident light is white. That will not be exactly the same as the color of the light coming directly from the interior of the sea, but the difference will not be great when there are many white clouds; it may be called the color by transmission.

E. O. Hulburt, 650 confining his observations to the "rim of the sea," the region bounded by lines of sight making angles of 0° and of 3°, respectively, with the surface of the sea-which region "comprises more than 99/100 of the total area of the sea within the view of the observer," if he is less than 1000 ft. above sca-level—has found that the rim of the sea when ruffled by breezes of 5 to 25 knots takes its color from the sky at an altitude of 30°, and that its brightness is about 0.25 of that of the sky near the horizon, the sky being clear. The light reflected from the sea is, of course, polarized. Hulburt has studied that polarization.

During his bathysphere descents into the open sea, W. Beebe 651 observed the light scattered horizontally by the sea. At a depth of 600 ft. he comments on the blueness of the illumination, which was still "brilliant" at 800 ft. When he looked out and down he "saw only the deepest, blackest blue imaginable, a color which in the spectrum [as seen in his small spectroscope | had vanished four hundred feet above, overlaid and superseded by violet." At 1400 ft. "the outside world was, however, a solid, blue-black world, one which seemed born of a single vibration-blue. blue, forever and forever blue," (See also Table 167). The persistence of the visual sensation of blue after the spectroscope shows that the light is violet is again referred to as puzzling. 652 E. O. Hulburt 653 has suggested that this sensation of blueness may arise from a fluorescence of the eye itself.

According to J. Y. Buchanan, 649 only three color-types are required for describing the color of the surface-water of the ocean. (1) Deep olivegreen, because of chlorophyll; observed near the edge of polar ice, and in

⁶⁵⁰ Hulburt, E. O., J. Opt. Soc. Amer., 24, 35-42 (1934).

⁶⁶¹ Beebe, W., Bull. N. Y. Zool. Soc., 33, 201-232 (1930).

cos Beehe, W., Science (N. S.), 80, 495-496 (1934). cos Hulburt, E. O., Idem, 81, 293-294 (1935).

certain other places. (2) Indigo. As one goes south from the Arctic the surface water assumes a pronounced indigo color, which persists until latitude 40° is passed. (3) Ultramarine. As one goes north from the equator the color persists as a pure and brilliant ultramarine until latitude 30° is passed. The passage between ultramarine and indigo is usually very rapid; the area of mixing is restricted.

48. OPTICAL ROTATORY POWER OF WATER

(For the Faraday effect and the Verdet constant, see Section 54.)

The natural optical rotatory power of water (the ability of water to rotate the plane of polarization of a beam of light passing through it) is believed to be precisely zero. That it is indeed exceedingly small is shown by such observations as those of F. Bates and R. F. Jackson 654 who observed for two 200-mm tubes an average rotation of only -0.0011° , the extreme single observations being -0.0043° and $+0.0003^{\circ}$, each occurring with the same tube. These values lie within the range of their experimental error.

A. A. Bless,⁶⁵⁵ whose error of setting was less than 0.02°, was unable to confirm F. Allison's conclusion ⁶⁵⁶ that water possesses a slight power to rotate the plane of polarization of light that passes through it while it is subjected to the action of x-rays.

49. DIELECTRIC PROPERTIES OF WATER

The dielectric properties of water are determined by the value of its dielectric constant (ϵ) and of its absorption index (κ) , and by the way these values vary with the conditions. Electrical conductivity is not a dielectric property of the substance, but it must be considered in the interpretation of experimental observations.

Symbols and Definitions.

Only the most frequently used symbols are listed here; others are defined where they appear.

 $(\epsilon, \epsilon', \epsilon'', \epsilon_0, \epsilon_1)$ Dielectric constant. The value of the dielectric constant (ϵ) , expressed in electrostatic units, is defined as the ratio of the mutual electrical capacity of a given pair of equipotential surfaces, fixed with reference to each other, when immersed in the dielectric to their capacity when immersed in a vacuum. If the polarizing of the dielectric is accompanied by a dissipation of energy, the apparent dielectric constant is a complex quantity, $\epsilon = \epsilon' - i\epsilon''$; $i \equiv \sqrt{-1}$. In such cases the real part (ϵ') is commonly called the dielectric constant. In certain cases, notably water, the dielectric constant (ϵ_0) at optical frequencies is much smaller than that (ϵ_1) under static conditions.

⁶⁵⁴ Bates, F., and Jackson, R. F., Bull. Bur. Stand., 13, 67-128 (S268) (1915).

⁶⁵⁵ Bless, A. A., Phys. Rev. (2), 33, 121-122 (A) (1929).

⁶⁶⁶ Allison, F., Idem, 31, 158-159 (A) (1928).

- (κ, e) Absorption index (κ) . If a plane, simple harmonic, electromagnetic wave is traveling through a dielectric in the direction of z, its amplitude at z=0 being A_0 and at z=z being A, then $A=A_0e^{-2\pi\kappa n e/\lambda_0}$, e (2.7183) being the base of the natural system of logarithms, and n and λ_0 having the values defined below. As the intensity (I) of such a wave varies as A^2 , $I=I_0e^{-4\pi\kappa n z/\lambda_0}$. Either of these two equivalent formulas may be used to specify the significance of κ .
- $(n, c, \lambda, \lambda_0)$ The index of refraction (n) is c/V where c (2.9979 \times 10¹⁰ cm/sec) is the phase velocity of the wave in a vacuum, and V is that in the dielectric. Also, $n = \lambda_0/\lambda$ where λ_0 is its wave-length in a vacuum and λ is that in the dielectric.
- (k',s) The conductivity (k') is, by definition, equal to the longitudinal electrical conductance of a cylinder of the material of unit length and unit cross-sectional area. It is the reciprocal of the volume resistivity. By definiton, $s \equiv k' \lambda_0 / \epsilon c$ when all quantities are expressed in the same system of units; say, the cgse. If, however, k' is expressed in the cgsm system, and ϵ in the cgse, then $s = ck' \lambda_0 / \epsilon$. Both expressions have been used, frequently without any specific statement about the units. Here, only the first will be used, the one in which all quantities are expressed in the same system.
- (ω, ν) A simple harmonic oscillation will be expressed either as $B_0 \sin \omega t$ or as the real part of $B_0 e^{i\omega t}$; $i \equiv \sqrt{-1}$. Then $\omega = 2\pi\nu$, where ν is the frequency of the oscillation.
- (ϕ, θ) The phase defect (ϕ) of a dielectric may be defined as follows. If a given pair of equipotential surfaces, fixed with reference to each other, distant from all others, and immersed in the dielectric, are subjected to a difference in potential defined by $V = V_0 \sin \omega t$, then the current will be $J = J_0 \cos (\omega t \phi)$, and the polarization of the dielectric will be $D = D_0 \sin (\omega t \theta)$. Both ϕ and θ depend upon the properties of the dielectric. They differ from zero only when the process of polarizing the dielectric is accompanied by a dissipation of energy, and the value of ϕ is given by the relation $\tan \phi = \epsilon''/\epsilon'$. If the dissipation arises solely from the process of polarizing the dielectric, k' being zero, then $\theta = \phi$; but if the dissipation arises solely from the conductivity, then $\theta = 0$, and $\tan \phi = 2s = 2k'\lambda_0/\epsilon c = 2k'/\epsilon v$.

Types of Dielectrics.

Two extreme types of dielectrics may be distinguished: (1) The ideal leaky dielectric, which is equivalent to an ideal, non-conducting and unabsorbing dielectric in parallel with a conductor of low conductivity. The combination exhibits both dispersion (variation of ϵ with λ_0) and absorption;

$$\epsilon = n^2(1-\kappa^2), \ \kappa = \frac{2s}{1+(1+4s)^{\frac{1}{2}}}, \ \tan \phi = 2s = 2k'\lambda_0/\epsilon c = 2k'/\epsilon \nu, \ \theta = 0.$$

Insofar as current-voltage relationships are concerned, the dielectric acts as if it were a non-conducting, absorbing dielectric with the dielectric constant $\epsilon_{\theta} = \epsilon - i4\pi k'/\omega = n^2(1 - i\kappa)^2 = n^2(1 - \kappa^2) - 2in^2\kappa$. This case will

not concern us further. Hereafter it will be assumed that either k' is zero or, if not, that effects arising from it have been independently eliminated.

(2) The non-conducting dielectric, either absorbing or non-absorbing, the last being but a limiting case. For such a dielectric, $\epsilon = n^2(1 - i\kappa)^2 =$ $n^2(1-\kappa^2) - 2in^2\kappa = \epsilon' - i\epsilon''$, tan $\phi = \epsilon''/\epsilon'$. As these expressions for ϵ are of identically the same form as those for the effective dielectric constant (ϵ_e) in the preceding case, $\epsilon''\omega/4\pi = \epsilon''\nu/2$ may be called the apparent conductivity of the dielectric (it is frequently called simply the conductivity, or more recently, the dipole conductivity, although k' is, by hypothesis, zero). For such dielectrics (k'=0), the dissipation of energy, arising solely from the act of changing the polarization, is intimately bound up with the true dielectric properties. The resulting absorption of energy is, therefore, commonly called dielectric absorption. (It was first described as "anomalous," to indicate that it did not arise from the conductivity as commonly measured.) It is in this sense that the term "dielectric absorption" is used in this compilation. It should not be confused with what Maxwell called electric absorption, which is merely one of the phenomena that accompany dielectric absorption.

Dipole Theory.

In the modern dipole theory of dielectrics, developed by Debye, the molecule is pictured as containing, in addition to the elastically bound electrons and ions of the earlier theories, a rigid or semirigid permanent electrical dipole firmly attached to the molecule, so that both move as a single unit. For certain types of molecules the moment of the dipole may be zero; for those substances the dipole theory adds nothing to the earlier ones. Thirty-five years ago M. Reinganum 657 suggested that certain molecules contained dipoles of constant moment, but whether the dipole could rotate without rotating the molecule itself was left an open question.

The presence of dipoles confers upon the dielectric two new types of polarization: (1) that produced by the aligning of the axes of the dipoles with the direction of the field, as a result of a reorientation of the molecule as a whole; and (2) that produced by the mutual angular displacement of the axes of adjacent dipoles that are elastically coupled, as two magnets might be, by their mutual attraction. This last is exactly similar to that associated with oscillators of other types, and needs no further consideration here, being completely covered by the well-known treatment of optical dispersion, together with P. Drude's extension of that to the case of great damping. 658

Of the first of these types of polarization, two subtypes need to be considered: (1) that characterized by a free reorientability (free rotation) of each molecule at every instant, and (2) that characterized by a restricted reorientability (restricted rotation), each molecule being elastically bound

⁶⁵⁷ Reinganum, M., Ann. d. Physik (4), 10, 334-353 (1903).
⁶⁵⁸ Drude, P., Z. physik. Chem, 23, 267-325 (1897) esp. Ann. d. Physik (Wied.), 64, 131-158 (1898).

to its neighbor or neighbors in such a way that the moment of the resultant dipole is not zero, or each molecule being sometimes bound and sometimes free. Until very recently only the first of these subtypes had been mathematically considered.

Free reorientability.—Debye has considered in detail the case of freely reorientable dipoles. His results have been published in numerous papers, and the basic treatment has been given in his book, "Polar Molecules," (1929), 658a from which much of the following has been derived, and to which reference will be made by means of the symbol PM followed by the number of the page.

In Debye's treatment it is assumed (a) that adjacent dipoles are not elastically bound to one another; (b) that each dipole plays the same part as every other; (c) that such a rotation of the molecule as attends the aligning of the axis of the dipole with the field is resisted by a torque of a viscous nature; (d) that the alignment of the axes is being continually disturbed by the thermal agitation of the molecules; and (e) that the Clausius-Mossotti expression for the molar polarizability of the dielectric, $P = (M/\rho)(\epsilon - 1)/(\epsilon + 2)$, is applicable. The last assumption implies that in the computation of the electric field at any point in the interior of the dielectric, the dielectric may be treated as a continuous medium, and the effect of a vanishingly small volume of the dielectric immediately surrounding the point may be ignored. As the theory deals with a molecular medium, the validity of this assumption has been questioned; and a steadily increasing amount of experimental data has forced the conclusion that this simple theory is certainly not quantitatively applicable to strongly dipolar pure liquids. But for years it was applied to them, and this must be borne in mind when studying the work of that period.

On the basis of these assumptions, Debye (PM, 27) derived, by statistical methods, expression (1) for the case of a constant field

$$\frac{P}{M} \equiv \left(\frac{\epsilon - 1}{\epsilon + 2}\right) \frac{1}{\rho} = \frac{4\pi N}{3M} \left[\alpha + \frac{\mu}{F} L\left(\frac{\mu F}{kT}\right)\right] \tag{1}$$

where
$$L(x) = \coth x - \frac{1}{x} = \frac{x}{3} \left(1 - \frac{x^2}{15} + \frac{2x^4}{315} - \dots \right)$$
. If only the first,

or the first two, terms in this expansion are retained, (1) becomes (2) or (3)

$$\frac{P}{M} = \left(\frac{\epsilon - 1}{\epsilon + 2}\right) \frac{1}{\rho} = \frac{4\pi N}{3M} \left[\alpha + \frac{\mu^2}{3kT}\right] = a + \frac{b}{T}$$
 (2)

$$\frac{P}{M} \equiv \left(\frac{\epsilon - 1}{\epsilon + 2}\right) \frac{1}{\rho} = \frac{4\pi N}{3M} \left[\alpha + \frac{\mu^2}{3kT} \left(1 - \frac{\mu^2 F^2}{15k^2 T^2}\right)\right] \equiv \alpha + \frac{b}{T} - \frac{f}{T^3}$$
 (3)

Here P is the molar polarizability of the substance (often called molar polarization), M molecular weight, ρ density, N Avogadro's number (6.061 \times 10²³ molecules per g-mole, k Boltzmann's molecular gas constant

 1.372×10^{-16} erg/°K per molecule), T °K the absolute temperature, F the strength of the internal field, α the polarizability of the molecule by elastic displacement of the electrons, and μ the moment of the dipole. The quantities α and μ refer to the individual molecule; α , β , and β are merely symbols to be used for brevity. It is obvious that in all cases the value of ϵ can be derived from P by means of the relation

$$\epsilon = (1 + 2P\rho/M)/(1 - P\rho/M) \tag{3a}$$

As the intensity of the field is increased, the dipole contribution to the polarization approaches a condition of saturation, and the corresponding term in the polarizability P begins to decrease. This is shown by the presence of the F^2 term in expression (3). By means of formula (3a) one can readily derive from (3) expression (4) in which ϵ_r is the dielectric constant in an external field of intensity X and ϵ_l is that in a weak field, it being remembered that on the Mossotti hypothesis the internal field is $F = X + 4\pi I/3$ and $\epsilon_l = X + 4\pi I/3$. I being the polarization of the medium; this gives us $F = X(\epsilon + 2)/3$. This ϵ is strictly ϵ_r , but as ϵ_r differs but little from ϵ_l the latter may be used in translating F into X in expression (3).

$$\epsilon_{\alpha} = \epsilon_{l} - \frac{4\pi}{45} \frac{N\rho\mu^{2}}{MkT} \left(\frac{\epsilon_{l}+2}{3}\right)^{4} \left(\frac{\mu X}{kT}\right)^{2} \tag{4}$$

or

$$\frac{\epsilon_{l} - \epsilon_{x}}{\epsilon_{l}} = \frac{4\pi}{45} \cdot \frac{N\rho\mu^{2}}{MkT\epsilon_{l}} \cdot \left(\frac{\epsilon_{l} + 2}{3}\right)^{4} \cdot \left(\frac{\mu X}{kT}\right)^{2} \tag{5}$$

Debye has used n to denote N_ρ/M , the number of molecules per unit of volume. In addition to this difference in notation, it will be noticed that the magnitude of the negative term in (4) is only a third as great as that of the one given by Debye (PM, 111); Debye's value for ϵ_r is less than that given by (4). The procedure that he followed (PM, 110) leads to an expression of the form $\epsilon_r = (1+2\delta)/(1-\delta)$, which may be expanded into $\epsilon_x = 1+3\delta+3\delta^2+\ldots$, since $\delta < 1$. If all powers of δ higher than the first are neglected, one obtains Debye's expression for ϵ_x , an expression that errs on the side of being too small. But δ may be, and in the case of water is, not much smaller than unity, in which case its higher powers cannot validly be neglected. This may be the explanation of the loss of the factor 1/3 from Debye's expression, which seems to be still current. 659

For water at 20 °C ($\epsilon_l = 80.4$ cgse, approximately) formula (5) reduces to

$$(\epsilon_l - \epsilon_x)/\epsilon_l X^2 = 0.0102_4 (10^{18} \,\mu)^4 \,\% \text{ per } (10 \text{ cgse field strength})^2$$
 (6)
= 0.00114 $(10^{18} \,\mu)^4 \,\% \text{ per } (\text{kilovolt/cm})^2$ (7)

For the H₂O molecule, $(10^{18} \,\mu)^4$ is approximately $11\frac{1}{4}$ cgse units (see p. 48), and $(\epsilon_l - \epsilon_x)/\epsilon_l X^2 = 0.0128\%$ per (kilovolt/cm)². Under the action of an electric field, molecules that are "elastically"

ess See Debye, P., Chem'l Rev., 19, 171-182 (1936).

aeolotropic tend to orient themselves so that the direction of maximum polarizability lies along the direction of the field. Consequently the presence of such molecules will give rise to effects that are strictly analogous to those produced by molecules containing fixed dipoles (c.f. PM, 109).

For a sinusoidal field of frequency v, Debye (PM, 90) finds

$$\frac{P}{M} \equiv \left(\frac{\epsilon - 1}{\epsilon + 2}\right) \frac{1}{\rho} = \frac{4\pi N}{M} \left(\alpha + \frac{\mu^2}{3kT} \cdot \frac{1}{1 + i\omega\tau}\right) \tag{8}$$

where $\omega = 2\pi v$ and τ is the relaxation time, which he defines as the time required for the polarization (not the polarizability P) to decrease to 1/eof its value after the constant inducing field is withdrawn. If the frictional torque resisting the turning of the molecule is $\zeta d\theta/dt$, $d\theta/dt$ being the angular velocity of the molecule, then $\tau = \zeta/2kT$ (PM, 94).

Drude-Debye relations.--At extremely high frequencies the massive molecules cannot follow the field, and formula (8) reduces to its first term. The value of the dielectric constant under those conditions will be denoted At very low frequencies (8) reduces to (2); that static value of the constant will be denoted by ϵ_1 . Then from (8) the following isothermal formulas may be derived, it being tacitly assumed, in (8) as well as in the following formulas, that after leaving the optical spectrum there is one, and only one, type of singularity in ϵ , and that that is of the type here covered.

There is nothing about these isothermal equations that is peculiar to the dipole theory 660; they may all be obtained from Drude's paper 600a as has been remarked by J. Malsch, 661 W. Ziegler, 661a and others. For this reason they are here called the Drude-Debye relations.

But the forms of the expressions given by Drude are not always the same as those given by Debye (PM, 90-94), and the time factor (a) that he used differs from Debye's τ by a factor that is a constant for any one Both forms, as well as varieties of each, are here given; as one form may, for certain purposes, be preferable to another.

For brevity, write $x \equiv \omega \tau (\epsilon_1 + 2)/(\epsilon_0 + 2) \equiv a\nu$ and $\eta^2 \equiv (\epsilon_1^2 + \epsilon_2^2)/(\epsilon_0^2 + \epsilon_2^2)$ $\epsilon_0^2 x^2$)/(1 + x^2); as usual, $\omega = 2\pi v$. Then the several quantities occurring in the expression $\epsilon = \epsilon' - i\epsilon'' = n^2(1 - i\kappa)^2$ may be computed by means of the following formulas:

$$\epsilon' = n^2(1 - \kappa^2) = (\epsilon_1 + \epsilon_0 x^2)/(1 + x^2) = \epsilon_0 + (\epsilon_1 - \epsilon_0)/(1 + x^2) = \epsilon_1 - (\epsilon_1 - \epsilon_0)x^2/(1 + x^2)$$
(9)

$$\epsilon'' = 2n^2\kappa = (\epsilon_1 - \epsilon_0)x/(1+x^2) = (\epsilon_1 - \epsilon_0)a\nu/(1+a^2\nu^2)$$
 (10)

$$2n^2 = \eta + \epsilon'; 2n^2\kappa^2 = \eta - \epsilon' \tag{11}$$

$$\tan \phi = \epsilon''/\epsilon' = 2\kappa/(1 - \kappa^2) = (\epsilon_1 - \epsilon_0)x/(\epsilon_1 + \epsilon_0 x^2) = [(\epsilon_1 - \epsilon')(\epsilon' - \epsilon_0)]^{\frac{1}{2}}/\epsilon'$$
(12)

⁶⁰⁰ Cf. Oplatka, G., Helv. Phys. Acta, 6, 198-209 (1933); Murphy, E. J., Trans. Electroch. Soc. (Amer.), 65, 133-142 (1934).

⁶⁰⁰a Drude, P., Ann. d. Physik (Wied.), 64, 131-158 (1898). 601 Malsch, J., Ann. d. Physik (5), 19, 707-720 (1934).

⁰⁸¹a Ziegler, W., Physik. Z., 35, 476-503 (1934).

$$\kappa = [(\eta - \epsilon')/(\eta + \epsilon')]^{\frac{1}{2}} = (\epsilon_1 - \epsilon_0)x/(\eta + \epsilon') (1 + x^2) = \tan (\phi/2)$$
(13)

From the expression defining x and a, we find

$$a = 2\pi\tau(\epsilon_1 + 2)/(\epsilon_0 + 2); \tau = a(\epsilon_0 + 2)/2\pi(\epsilon_1 + 2)$$
 (14)

For deriving the value of a from the observed values of ϵ' and $\epsilon'' \equiv 2n^2\kappa$, Drude advises the use of one or another of the following formulas:

$$a^{2} = (\epsilon_{1} - \epsilon')/(\epsilon' - \epsilon_{0})\nu^{2}; a = (\epsilon_{1} - \epsilon')/2n^{2}\kappa\nu;$$

$$a = 2n^{2}\kappa/(\epsilon' - \epsilon_{0})\nu$$
 (15a, 15b, 15c)

From (10) one finds

$$2x = A - (A^2 - 4)^{\frac{1}{4}}, \text{ where } A \equiv (\epsilon_1 - \epsilon_0)/2n^2\kappa$$
 (16)

from which τ can be found from the relation:

$$\tau = x(\epsilon_0 + 2)/2\pi(\epsilon_1 + 2)\nu \tag{17}$$

It will be noticed that when the frequency is such (v_8) that x = 1, then $a=1/\nu_{e}, \ \tau=(\epsilon_{0}+2)/(\epsilon_{1}+2)2\pi\nu_{e}, \ \text{and} \ \epsilon'=(\epsilon_{1}+\epsilon_{0})/2.$ This frequency, at which ϵ' is midway between ϵ_1 and ϵ_0 , may be called the transition frequency. It is characteristic of the substance, fixing the values of the coefficients a and τ , which have throughout this treatment been regarded as independent of the value of v. The corresponding wave-length in a vacuum $(\lambda_8 = c/\nu_8)$ may be called the transition wave-length; $\lambda_8 = ca = \lambda_0 a\nu = \lambda_0 x$, where $\lambda_0 = c/\nu$. The quantity $c\tau$ also defines a characteristic wave-length; one that is much smaller than the transition wave-length. Several other characteristic wave-lengths may be defined (see following table). A reader must be on the alert, for authors do not always state clearly which of the several characteristic wave-lengths is being considered. As they all serve to fix the region in which the dielectric constant changes rapidly, each may be called a transition wave-length, but here the term will be restricted to the one for which ϵ' is midway between ϵ_1 and ϵ_0 . This is the one most frequently designated by λ_s and called by Germans "Sprungwellenlange."

In the following table, the second column contains the values of x_0^2 or $a^2\nu_0^2$ ($x\equiv a\nu$) corresponding to the criterion specified in the first column

("relaxation" means the time (τ) required for the polarization to become reduced to 1/e of its value, the impressed field being zero), and in the last column are given the values for water at 20 °C of each of these characteristic wave-lengths in terms of transition wave-length (λ_e) and of λ_{τ} as defined by the relation $\lambda_{\tau} = c\tau$.

If in (15b) and (15c) a is replaced by its equivalent λ_s/c and ν by c/λ_0 , then one readily obtains

$$\epsilon' = \epsilon_1 - (2n^2\kappa/\lambda_0)\lambda_s$$
 and $\epsilon' = \epsilon_0 + 2n^2\kappa\lambda_0/\lambda_s$ (18a, 18b)

That is, if there is but one such jump as we are considering and there is no absorption band beyond the optical spectrum, then ϵ' will be linear in both $2n^2\kappa/\lambda_0$ and in $2n^2\kappa\lambda_0$, and the values of ϵ_1 and ϵ_0 will be given by the intercept of these lines on the axis of ϵ' .

Restricted reorientability.—If as the result of their mutual action the dipoles of adjacent molecules are elastically coupled, as two magnets may be, then their combined moment will be less than the sum of the single, separate moments, and may in the limit be zero. Such coupling may be either permanent or temporary. All of which will tend to reduce the magnitude of that portion of the dielectric constant that is contributed by the dipoles, and of the variation of that portion with the strength of the field.

Debye 662 has recently considered this problem, and has derived formulas for the case in which each molecule is so elastically bound to a direction fixed in space that the potential energy of the molecule contains a term of the form $-E\cos\theta$ where θ is the angle that the axis of the dipole makes with the fixed direction. That fixed direction varies from molecule to molecule, and for each molecule it is continually changing as a result of the thermal agitation of the molecules. The compiler has not yet been able to perceive clearly the physical significance of these assumptions. Debye concludes that E is very great, and that the effect of restraints of the kind just specified is to multiply the b and the f of formula (3) by 2C and $3C^4$ respectively, C standing for E/kT. As the f of formula (3) is proportional to the last term in (4), the restrictions here imposed require that also to be multiplied by $3C^4$.

For other suggested explanations of the failure of liquids to conform to Debye's formulas for freely reorientable dipole molecules, and for criticisms of Debye's treatment of restricted reorientability, see M. Forró, 663 J. Frenkel, 664 G. Hettner, 665 J. Malsch, 666 L. Onsager, 667 and A. Piekara. 668

Dielectric Constant of Water.

Miscellanea.—The degree of accord between various theories and the experimentally determined values for various substances has been discussed by O. Blüh.⁶⁶⁹ The data for water, ice, and steam are considered in detail. The article concludes with a bibliography of 172 entries. More recently,

⁶⁶² Debye, P., Acad. roy. Bela Bull. Cl. Sci. (5), 21, 166-174 (1935); Physik. Z., 36, 100-101, 193-194 (1935). Chem'l Rev., 19, 171-182 (1936).

ees Forró, M., Z. Physik, 47, 430-445 (1928).

⁶⁰⁴ Frenkel, J., Acta. Physiochim. URSS, 4, 341-356 (1936).

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one Malsch, J., Ann. d. Physik (5), 29, 48-60 (1937).

⁶⁰⁷ Onsager, L., J. Am. Chem. Soc., 58, 1486-1493 (1936).

⁶⁰⁸ Piekara, A., Acta Phys. Polon., 6, 130-143 (1937).

- W. Ziegler ⁶⁷⁰ has done the same, but with special reference to the dispersion and absorption of electric waves. He gives a bibliography of 159 entries. For discussions of the difficulties inherent in the use of waves along wires immersed in the dielectric, see E. Frankenberger. ⁶⁷¹ Among other things he concludes that changes in the surface of the wire may introduce systematic errors of several tenths of a per cent in *n*, the index of refraction.
- P. Drude 672 has concluded that dissolved air produces no observable effect upon the value found for ϵ . Both he and A. Deubner 673 comment on the fact that the successive individual determinations of ϵ for a given sample of water are more concordant than are the determinations for different samples. Although this suggests that nominally identical specimens of water have different values of ϵ Drude dismisses such a suggestion as being extremely improbable.
- G. Jacoby 674 has derived an expression relating the dielectric constant to the forces binding the atoms in the molecule. From this he concludes that one of the hydrogens in H_2O is bound more loosely than the other.

The dielectric constant of a film of water 2 microns (0.0002 cm) thick is the same as that of water in bulk.⁶⁷⁵ The following papers touch upon the dielectric constant of water, but are not mentioned elsewhere in this section: James Dewar and J. A. Fleming ⁶⁷⁶ discuss the data obtained for water prior to 1897, and ⁶⁷⁷ give a single very low value for 1 °C. R. Fürth, ⁶⁷⁸ M. Jezewski, ⁶⁷⁹ H. Joachim, ⁶⁸⁰ Y. Matsuike, ⁶⁸¹ E. B. Rosa, ⁶⁸² J. F. Smale, ⁶⁸³ S. Tereschin, ⁶⁸⁴ B. B. Turner, ⁶⁸⁵ and G. U. Yule, ⁶⁸⁶ each reports a single determination, generally incidental to another problem or for the purpose of testing a proposed procedure. W. Nernst ⁶⁸⁷ reports a few preliminary determinations.

Saturation.—On Debye's simple theory of freely reorientable dipoles, the dielectric constant of a dipole substance should exhibit "saturation" effects in intense fields, the dielectric constant (ϵ_i) in a strong field of intensity X being less than that (ϵ_i) in a weak field, by the relative amount

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one Blüh, O., Phy. Z., 27, 226-267 (1926).
670 Ziegler, W., Phys. Z., 35, 476-503 (1934).
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672 Drude, P., Ann. d. Physik (Wied.), 59, 17-62 (1896).
<sup>678</sup> Deubner, A., Ann. d. Physik (4), 84, 429-456 (1927) = Diss., Freiburg. <sup>674</sup> Jacoby, G., Ann. d. Physik (4), 72, 153-160 (1923).
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676 Dewar, James, and Fleming, J. A., Proc. Roy. Soc. (London) (A), 61, 2-18 (1897).
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688 Smale, J. F., Ann. d. Physik (Wied.), 57, 215-222 (1896).
684 Tereschin, S., Idem, 36, 792-804 (1889).
ees Turner, B. B., Z. physik. chem., 35, 385-430 (1900).
ene Yule, G. U., Ann. d. Physik (Wied.), 50, 742-751 (1893).
687 Nernst, W., Z. physik. chem., 14, 622-663 (1894).
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 $(\epsilon_l - \epsilon_w)/\epsilon_l = AX^2$, A being a factor depending upon the substance and its temperature (see eq. 5). For water, his expression, when corrected (p. 354), leads to the computed value $A_o = 0.0128$ per cent per (kilovolt/cm)².

But J. Malsch ⁶⁸⁸ has found for water A = 0.000011 per cent per (kilovolt/cm)², which is only 1/1000 of A_c . Debye's theory for restricted reorientability (p. 357) can be made to account for such a factor.

Variation with Frequency.—From a study of all pertinent data available in 1924 (a few later data were studied and placed in the report as it was being prepared for the printer), H. L. Curtis 689 concluded that, for ν not exceeding 100 megacycles/sec, the dielectric constant (ϵ) of water at a pressure of one atmosphere is independent of the frequency, and is about 81.2 at 17 °C. At higher frequencies, beginning near $\nu = 600$ megacycles/sec, ϵ steadily decreases as shown in Figure 5.

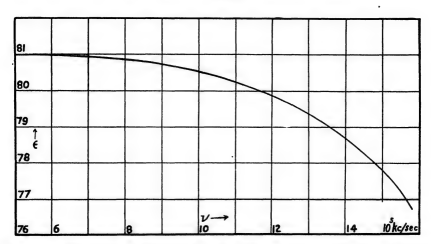


FIGURE 5. Dielectric Constant of Water at 17 °C; Variation with the Frequency.

[From compilation by H. L. Curtis and F. M Defandorf, Int. Crit. Tables, 6, 77 (1929).]

Unit of ν (frequency) is 100 megacycles per second; of ε (dielectric constant) is 1 cgse.

The most exact determinations of the dielectric constant of water now available at the lower frequencies (under 100 megacycles/sec) are probably those of J. Wyman, Jr. 690 and of F. H. Drake, G. W. Pierce, and M. T. Dow. 691 They found, respectively, 81.47 and 81.54 at 17 °C. Their mean, 81.50, is used in Table 174 and is probably as close an estimate of the correct value as can be obtained from the data available. It differs by only 0.7 per cent from the mean (80.9) of the entries in the first section of Table 172, the two plainly abnormal values (88 and 73) being ignored.

⁶⁸⁸ Malsch, J., Physik. Z., 29, 770-771 (1928); 30, 837-839 (1929).

⁶⁸⁹ Curtis, H. L., Int. Crit. Tables, 6, 77, 78 (1929).

Wyman, J., Jr., Phys. Rev. (2), 35, 623-634 (1930).
 Drake, F. H., Pierce, G. W., and Dow, M. T., Idem, 35, 613-622 (1930).

The existence of bands of anomalous dispersion at frequencies below 1000 megacycles/sec seems to be generally discredited at present. Although such bands have been reported by A. R. Colley,⁶⁹² K. Iwanow,⁶⁹³ and R. Weichmann,⁶⁹⁴ others have been unable to confirm their existence, and at least some of them may have been caused by an impurity, probably glass, dissolved in the water (see G. Mie,⁶⁹⁵ E. Frankenberger ⁶⁹⁶). Likewise the report by A. Bramley ⁶⁹⁷ of a region of selective absorption, and hence, presumably, of anomalous dispersion, was later found by himself to be incorrect.⁶⁹⁸ The regions of low frequency in which water has been especially investigated for the existence of absorption bands are listed in Table 173.

Table 172.—Dielectric Constant of Water at 17 °C

The best available value for frequencies below 600 megacycles per second is 81.50 cgse unit. It seems improbable that there are bands of absorption at frequencies below 1000 megacycles/sec, although such bands have been occasionally reported (see text and Table 173).

When an observer has concluded that his observations indicate that ϵ or n is the same for each of the frequencies that he has used, the mean of all his values has been tabulated together with the number (No.) of frequencies studied. If he has actually determined n, the value of n^2 is given, and ϵ (actually ϵ' , the real part of ϵ) is also given if the author has either computed it himself or given data that enables one to determine it. Under Osc. or Os. is an indication of the damping of the oscillations used: d = damped, d = slightly damped, u = undamped. Below 1000 megacycles/sec the absorption of the water is so slight that n^2 is essentially the same as ϵ .

Unit of $\lambda_0 = 1$ cm, of $\nu = 1$	megacycle/sec, of e= :	1 cgse unit.	Temp. = 17 °C
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λ ₀	v		n²	No.	Osc.	Ref.
$5.0(10^8)$	60(10-6)	81.08	**		u	Ca
5.0(104)	0.6	81.2			u	Ko
3.5(104)	0.86	81.9			u	Ку
2.0(104)	1.5	88.1			u	PH
1.1(104)	2.7	82.5			u	As
8109-368	3.70-81	81.5°		5	u	Wy
2550-392	12-76	81.5	$\mathbf{w}^{\mathbf{d}}$	2	u	$\vec{\mathbf{D}}\vec{\mathbf{D}}$
1110-309	27-97		79.4	3	ã	\overline{CZ}
700-332	43-90		81	7	u	MJ
600-310	50-97		80.5	2	d	Co
444-242	68-124	73	Wd	4	u	Sa
345-230	87-130		81.07	7	ā	Ã1
321-36.7	93-818		80.68	7	u	He
290-230	103-130		81.00	4	u	No
276-124	109-242		80.0	2	น	So

^{∞∞} Colley, A. R., Phys. Z., 10, 471-480 (1909) \leftarrow J. Russ. Fis.-Chim. Obsc. (Phys.), 39, 210-233 (1907).

⁶⁰⁸ Iwanow, K., Ann. d. Physik (4), 65, 481-506 (1921).

^{66, 501-545 (1921) →} Phys. Z., 22, 535-544 (1921).

es Mie, G., Phys. Z., 27, 792-795 (1926).

con Frankenberger, E., Ann. d. Physik (4), 82, 394-412 (1917); (5), 1, 948-962 (1929).

⁶⁰⁷ Bramley, A., J. Franklin Inst., 206, 151-157 (1928); 207, 315-321 (1929) → Phys. Rev. (2), 33, 279 (A), 640 (A) (1929).

oos Bramley, A., Phys. Rev. (2), 34, 1061 (L) (1929); J. Opt. Soc. Amer., 21, 148 (A) (1931).

				Tab	le 1	72(Conti	писа	1)					
	X ₀ 271-268 266 250-60 200-25.8 200 75 73.4-23.8 68.4-55.2 63.7-59.5 63-49 56.6-52.2 56.7-28.4 49.9-39.4 45.6 35.0-31.0	2	113 120 150 400 400 409 438 471 476 530 601 658	1-500 1-1160 1-1260 1-542 1-504 1-612 1-574 1-1056 1-762		•	•		80.82 81.5 80.4 81.8 79.7 80.2 81.7 81.1 80.3 81.3 79.8 80.5 80.1 81.0 81.1 81.1		No. 12 14 8 73 20 5 34	u d d d 3 d u d u d		Ref. De Ho Ho Tr' Dr' Fr' Cy GA Fr' AOP RD Rp
	Ref.a→ Osc.→ v/1000	Rp d	Go	Kn u	Se u n²	Ec d	Te ^f	La d	n ²		Ref.a	Kn u	Ec d	Te d
25.8 24.43 24.06 23.82 23.06	1.16 1.23 1.25 1.26 1.30	79.1 78.7 78.0	98.8		,				82.2	ď	Ma			
21.59 20.55 20.44 19.0 17.76	1.39 1.46 1.47 1.57 1.69	77.8 77.8	91.4	77.4	80							77.3		
17.42 17.02 16.83 15.29 14.48	1.72 1.76 1.78 1.96 2.06		87.4 87.6	77.4 77.4 77.4								77.2 71.1 77.1		
14.0 13.6	2.14 2.21				84 82				77.0		4.0			
13.45 12.65 12.6	2.23 2.37 2.38		83.2		80				77.8	u	AO			
11.80 11.48	2.54 2.62		81.0	77.4								77.0		
11.11 10.20	2.69 2.94			77.4 77.4								77.0 76.9		
9.85 9.55 8.80 8.53	3.05 3.16 3.43 3.51		74.0	77.4 77.4 77.4		78.3						76.9 76.9 76.8	77.9	
8.05 7.1	3.73 4.22			77.4					80.2		Mz	76.7		
6.48 6.2 5.7	4.63 4.84			77.4					86.5		Mz	76.3		
5.7 5 4	5.26 6.0					78.8			77.80	đ	Co		77.6	
3.7 2.7 1.8	7.5 8.1 11.1 15.6					64.4	72.5 45.4		76.1	đ	El		61.4	67.4 40.0

Table 172.—(Continued)

O	ef."→	Rp d	Go u	Kn u	Se Ec u d	Te/ d	La d	Misc. — Misc. — Os. Ref.a	Kn u	Ec d	Te d
λ ₀ 1.75	ν/1000 17.2			n²	63.8			n ² Os. Ref. ^a		56.5	
1.5	20.0 27.3					45.0 40.4					41.6 38.4
0.84 0.8	35.7 37.5					33.4	82.7				31.2
$0.6 \\ 0.42$	50.0 71.4					29.6	89.7				27.9
0.4 0.005	75.0 6000			(Optica	1)		91.4	1.85			

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- Wyman, J., Jr., Phys. Rev. (2), 35, 623-634 (1930).
- ⁶ From the constant of an electrometer with its plates immersed in water.
- From the frequency of an oscillator immersed in water.
- ^d Both DD and Sa measured n, but reported only the values of e as derived from the n's by correcting for the electrical conductivity.
- * Rukop determined n for 51 wave-lengths well distributed over the range $\lambda_0 = 63.7$ to 20.6 cm, and concluded that n is constant over the ranges 63.7 to 59.5, 49.9-39.4, and 35.0 to 31.0 cm. Between those regions and for $\lambda_0 < 31$ cm his observations were less uniform, and he thought there might be anomalous dispersion; but he could not find the sharp bands reported by Colley, and for which he was searching.
 - Tear measured the reflectivity and the absorption, and computed n from them.
- ^o Cole measured the reflectivity for $\lambda_0 = 5$ cm, and computed n from that on the assumption that there was no absorption.

Table 173.—Absorption Bands in the Electrical Spectrum of Water

It seems probable that water has no absorption bands at frequencies below 1000 megacycles/sec,600 but bands have been reported in certain of the regions here tabulated, all of which have been carefully searched for bands; the number of bands reported in each case is stated in column 3. The range 250 to 450 megacycles/sec should be reëxamined. λ_0 = wavelength in air, $\nu =$ frequency.

	Unit of $\nu = 100$ megacycle	es/sec; of $\lambda_0 = 1$ cm	
ν	λο	Bands	Ref.a
0.8 to 1.0	375 to 300	2 sets	\mathbf{Br}^{b}
0.84 to 1.06	260 to 280	None	AN
1.0 to 1.4	300 to 220	None	No
1.0 to 1.4	300 to 220	None	A 1
2.4 to 2.5	124 to 120	None	Iw^b
2.5 to 4.5	120 to 67	Many	Iw^b
4.5 to 5.0	67 to 60	None	Iw^b , Cy'
4.7 to 14.6	64 to 20	3	Rp
5.3 to 8.4	56 to 36	Many	Ċy''
5.4 to 10.7	56 to 28	None	AO
5.6, 7.1, 7.4	54, 42, 32	3	We^b
5.3 to 5.9	56 to 52	None	Mieb, Fr'
4.1 to 8.1	73 to 37	None	Fr''
15 to 60	20 to 5	None	Kn
150, 430, 1500	2.0, 0.7, 0.2	3	Te^{b}

a References:

- Al AN
- See Table 172, references, first of Al.
 See Table 172, references, second of Al.
 See Table 172, references.
 Bramley, A. ar Reported selective absorption; hence anomalous dispersion may be inferred. He concluded later that the observed effects were spurious, at least in AO Br

- Fr' Fr"
- interred. The concluded lates that the part 108 (olley, A. R., see Table 172, Cy. (olley, A. R., Physik Z., 10, 471-480 (1909) \leftarrow J. Russ. Fis.-Chim. Obs. (Phys.), 39, 210-233 (1907). Frankenberger, E., Ann. d. Physik (4), 82, 394-412 (1927). Idem (5), 1, 948-962 (1929). Iwanow, K., Ann. d. Physik (4), 65, 481-506 (1921). He calls the value of $2\pi\nu/10$ frequencies; in this table the values of ν itself are given, as computed from his values of $\lambda_0/2$.
- Kn
- requencies; in this table the values of ν itself are given, as computed from his values of $\lambda_0/2$. Knerr, II. W., Phys. Rev. (2), 52, 1054-1067 (1937). Mie, G. 605 He concluded from the observations of Fr' that the bands reported by Mie We are spurious.
- No
- Rp
- We are spiritude. Novosilzew, N., see Table 172, references. Rukop, II., see Table 172, references. Tear, J. D., Phys. Rev. (2), 21, 611-622 (1923). He reported selective absorption at these frequencies; hence anomalous dispersion may be inferred. Weichmann, R., Ann. d. Physik (4), 66, 501-545 (1921) \rightarrow Physik. Z., 22, 535-544 (1921). Mie concluded that these bands are spurious.

Variation with Temperature.—(See also pp. 353-355). At the time of the compilation by H. L. Curtis and F. M. Defandorf 700 the available data on the variation of the dielectric constant of water with the temperature, especially at the higher temperatures, were far from satisfactory. The best that could be done was to give the linear equation $\epsilon = 80 - 0.4(t - 20)$, equivalent to $\epsilon = 81.2 [1 - 0.005(t - 17)]$, the coefficient of t being restricted to a single significant figure. It was known that the dielectric

^b See note accompanying the corresponding reference.

⁶⁰⁰ Malsch, J., Ann. d. Physik (5), 19, 707-720 (1934).

⁷⁰⁰ Curtis, H. L., and Defandorf, F. M., Int. Crit. Tables, 6, 74-81 (1929).

constant has no maximum at 4 °C, but the observations of F. Ratz ⁷⁰¹ indicated that there might be a very indistinct maximum between 0 °C and 1 °C.

Since then, both L. Kockel ⁷⁰² and J. Wyman, Jr.⁷⁰³ have published consistent data for the entire range, 0 °C to 100 °C; F. H. Drake, G. W. Pierce and M. T. Dow ⁷⁰⁴ for the range 10 °C to 60 °C; and R. T. Lattey, O. Gatty, and W. G. Davies ⁷⁰⁵ two short series in the range, 14 °C to 18 °C. All of these are closely represented by formula (18c)

$$\epsilon = \epsilon_{17} \left[1 - 4.696 \left(\frac{t - 17}{1000} \right) + 10.2 \left(\frac{t - 17}{1000} \right)^2 \right]$$
 (18c)

Values which appear to be of a much lower accuracy have been published by A. C. Cuthbertson and O. Maass ⁷⁰⁶ for 0, 15, 25, 50, and 75 °C; and E. P. Linton and O. Maass ⁷⁰⁷ have published a formula said to represent their own (unpublished) observations over the range 0 °C to 50 °C. This formula does not agree satisfactorily with the other data and, owing to the meagerness of the information published, cannot be critically appraised except by such comparison.

Formula (18c) also represents quite satisfactorily the data published by F. Ratz, those over an 8.5° range that are summarized by A. R. Colley 706 by means of an equation, those from 0° to 50 °C defined by a formula pro(Continued on Page 366)

Table 174.—Variation of the Dielectric Constant of Water with the Temperature

The several values are compared with those of (ϵ_f) defined by the preferred formula (18c), which is used in the following form:

$$\left(\frac{\epsilon}{\epsilon_{17}}\right)_f = 1 - 4.696 \left(\frac{t - 17}{1000}\right) + 10.2 \left(\frac{t - 17}{1000}\right)^2; \ \epsilon_f = 81.50 \left(\frac{\epsilon}{\epsilon_{17}}\right)_f$$

the value 81.50 being the mean of the ϵ_{17} values obtained by Wyman and by Drake, Pierce, and Dow (see p. 359).

Values of ϵ_f and of (ϵ/ϵ_{17}) , are tabulated, and the values reported by the several observers are indicated by means of the quantities Δ and δ , which are defined by the relations $(\epsilon/\epsilon_{17}) = (\epsilon/\epsilon_{17})_f + \Delta$ and $\epsilon_{obs} = \epsilon_f (1 + \delta)$. At the bottom of each column of Δ is given the value of ϵ_{17} used by the observer in computing his smoothed values. For example, Wyman's $\epsilon_{17} = 81.47$, his value of ϵ/ϵ_{17} at 0 °C is $1.082_8 - 0.003 = 1.080$; consequently his ϵ_0 is 81.47(1.080) = 87.99; by means of his formula (see text) we find $\epsilon_0 = 78.54(1.1205) = 88.00$.

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⁷⁰⁸ Cuthbertson, A. C., and Maass, O., J. Am. Chem. Soc., 52, 483-489 (1930).

⁷⁰⁷ Linton, E. P., and Maass, O., Idem, 54, 1863-1865 (1932).

Table 174—(Continued)

It will be noticed that, except for fixed percentile errors, most of the short sets of older data given in the last sections of the table agree quite satisfactorily with the values (ϵ_f) defined by formula (18c).

Unit of $\epsilon = 1$ cgse unit; of $\Delta = 0.001$; of $\delta = 0.1\%$. Temp. = t °C.

		Unit of	e=1 0	gse un	t; of	$\Delta = 0.00$	$i; oi \delta =$	0.1%.	Temp.	∍t°C.			
References	2 →	Ko	Ko	Wy	DD_i		Су	LM	Dr	ICT.		Ratzd	
t er	(e/e17) f	Obs.				Co:	mputed				t	Observ (e/e17)	
0 88.25	1.0828	+2	0	-3		-1		+5	1.2			1.082	
10 84.22	1.0334	+4	ŏ	-1	0	-1	+1	0	+2 +1	T	0.1 0.33 0.75	1.082	-3
20 80.35	0.9859	-6	0	-100	0	0	+1 -1	+1	Ô	ŤÔ	0.75	1.081	-2
25 78.49	0.963_{1}	_	0	+1 +1	. 0	+1 +1	-2	+1 +2	-2	- 2	0.80	1.078	+2
30 76.67	0.9407	-3	0	+1	+1	+1	-4	+4	-3		1.03	1.077	
40 73.14	0.8974	-5	+1	+2	+3	+2		+9 +12	-5 -8	- 10	1.57	1.074	0
50 69.78 60 66.60	0.856_{2} 0.817_{1}	$\frac{-3}{-5}$	0	+2 +2 +2 +2	+3 +6 +9	T2		+12	-12	- 18	1.80	1.074	$-1 \\ +1$
70 63.57	0.7809	-2	ŏ	+2	, ,	+2			-16	- 40	2.20 3.04	1.067	T Ô
80 60.71	0.745_{0}	+1	-1	U		+2 +2 +3 +2 +3				55	3.43	1.065	0
90 58.02	0.7122	+2	-2	-1		+2					4.00	1.062	
100 55.58	0.6810	+2	-4	-2		+1				-89	4.16	1.062	0 -2
€17 81.50 E	81.50		81.2	31.47	81.5	4 81.19	80.26	82.0	81.65	81.2	€17	=81.2	24
Lattey,	Gatty,	and :	Davie	s (Tv	vo s	eries, 1	and 2	2) LD4					
• ,	• ,	(1)	(2)	•		,	(1)	(2)				(1)	(2)
t	es c	δ		t		ef		8	t		ef ,		3 —
14.0 82	2.92	-4.2		33.9)	75.26		+0.3	61.15		24		+9.1
	2.04		+2.2			73.32	-3.4	, 0.0	63.0	65		- 3.5	1 7.1
		-1.4	,	42.0		72.44	0.1	+1.4	72.4	62.		0.5	-0.2
	9.81		-1.4			71.11	+0.5	,	74.5			-1.3	0.2
	8.04		-4.2			68.97 -			80.15		.68	1.0	-1.8
		- 6.8	1,2	54.2		68.41	10.2	+1.2	81.3			-3.6	-1.0
			. 2. 6			2.7; Se	mion 2		0.10	00	.00	0.0	
			10: 3				ries 2,						
	ef	δ		t		ef		δ		t	€f		δ
\mathbf{D}_{0}	evoto*				3.5	77.21		3.9		7.2	77.7		- 86
3.8 80	6.70	-3.5	5		0.6	76.44		7.8		1.7	76.1	-	- 99
	5.92	-4.3	3		3.8	75.30			35	5.3	74.8		103 '
	5.25	-4.9)		5.3	74.42		1.6					
	5.09	-1.	l	39	9.2	73.42	-11	1.6		M	[ean	-	- 96
	4.30	-3.8											
	3.67	-3.				Tezews	kia				Coolid	gea	
	3.12	-4.				5020			3	3.5	86.8	_	- 1.1
	2.27	-3.0			2.3	87.32	+:	39		3.6	82.8		-1.2
	1.73	-4.		10	5.5	81.69	+	76		9.0	80.7		-2.5
	1.69	- 5.9		30	0.1	76.63	+ '	76		1.7	78.6		0.0
	1.46	-5.4		4	7.2	70.70	+	68		9.0	73.5	_	-1.3
	1.23	-3.			5.9	64.49			0.				
	0.93	-5.3			2.	57.50	+:	33		11/	[ean	_	-1.2
	0.85	8.0			9.5	55.62		32		44.	LOGII		
		-6.				00.0-				H	[errwa	gena	
	0.63 0.51	- 5.0			7	Mean	+	53		4.70	86.34	.,	- 9.8
					•	120011	•			9.85	84.28		- 8.9
	0.04	-5.								2.75	83.14		- 8.4
	9.71	-4.9				Cohr	1ª						
	8.48	- 8.0			0.5	04.4		07		4.65	82.40		- 8.6
	8.42	-9.8			9.5	84.4		97		6.35	81.75		-8.7
	7.86	-8.			0.5	84.0	- 1		20	0.75	80,07		-6.4
28.3 7	7.28	– 10. 2	2		6.8	81.6		99		3.	r		0.5
				19	9.8	80.4	-	96		IV	Iean	•	- 8.5
" Refere													
Cohn,	E.709												
Coolid	dge, W.	. D.710			180	•							

Conling, W. D.⁷¹⁰
Cy Colley, A. R. See Table 172, references.
DD Drake, F. H., Pierce, G. W., and Dow, M. T.⁷⁰⁴
Devoto, G. See Table 172, references.

Table 174—(Continued)

Table 174—(Conlinued)

Dr Drude, P. 672

Heerwagen, F., Ann. d. Physik (Wied.), 49, 272-280 (1893).

ICT Curtis, H. L., and Defandorf, F. M., Int. Crit. Tables, 6, 74-81 (1929).

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Ko Kockel, L. 702

I.D Lattey, R. T., Gatty, O., and Davies, W. G. 705

Linton, E. P., and Maass, O., J. Am. Chem. Soc., 54, 1863-1865 (1932).

Ratz, F., Z. physik. Chem., 19, 94-112 (1896).

Wy Wyman, J., Jr. 708

 $^{\circ}$ The progressive increase in ϵ as t increases above 30 $^{\circ}$ C, as well as the abnormally small values (not here recorded) found below 10 °C, are probably the result of errors in determining the temperature.

As the coefficient of t is given to only one significant figure, even the first figure

of the number is uncertain.

^d Ratz gives values for t = 4.43, 5.37, 5.47, 10, 20, and 30 °C also; for which, $\Delta = -2, +1, 0, -2, -18, \text{ and } -5, \text{ respectively.}$

Table 175.—Variation of the Polarizability of Water with the Temperature

The molar polarizability of a dielectric is $P = (M/\rho)(\epsilon - 1)/(\epsilon + 2)$, and Debye's theory for freely reorientable dipole molecules leads to the relation $P/M \equiv (\epsilon - 1)/(\epsilon + 2)\rho = a + b/T$ (eq. 2) where $b = 4\pi N \mu^2/\epsilon$ 9MkT; hence $PT/M \equiv (\epsilon - 1)T/(\epsilon + 2)\rho = b + aT$. For water, the reorientability of the molecules is restricted, and PT/M is not linear in T; it is however closely given by the relation

 $PT/M \equiv (\epsilon - 1)T/(\epsilon + 2)\rho = 107.13 + 0.2262T + 0.00127550T^2$ where $T \circ K$ is the absolute temperature. The calculated values given below are those defined by this formula; the observed ones are determined from the indicated values of ϵ and ρ , the former being the values of ϵ_f given in Table 174. If b = 107.13, $10^{19} \mu = 5.59$ cgse units per gfw of H₂O (see p. 176).

Unit of ϵ -1 cgse; of ρ -1 g/cm³. Temp.= t° C.

				P7	C/M	
	T	e	ρ	Obs	Calc	o-c
0	273.1	88.25	0.9999	264.03	263.87	+0.16
10	283.1	84.22	0.9997	273.33	273.34	-0.01
20	293.1	80.36	0.9982	282.92	282.94	-0.02
25	298.1	78.49	0.9971	287.81	287.84	-0.03
30	303.1	76.67	0.9957	292.83	292.80	+0.03
40	313.1	73.15	0.9922	302.98	302.92	+0.06
50	323.1	69.79	0.9881	313.30	313.30	0.00
60	333.1	66.60	0.9832	323.93	323.91	+0.02
70	343.1	63.57	0.9778	334.83	334.80	+0.03
80	353.1	60.71	0.9718	345.92	345.94	-0.02
90	363.1	58.02	0.9653	357.32	357.33	-0.01
100	373.1	55.58	0.9584	369.00	368.98	+0.02

(Continued from Page 364)

posed by P. Drude, 672 the values obtained by E. Cohn, 709 by W. D. Coolidge,⁷¹⁰ and by F. Heerwagen.⁷¹¹ See Table 174; formulas are given below.

⁷⁰⁸ Colley, A. R., Physik. Z., 10, 329-340 (1909).

⁷⁰⁰ Cohn, E., Ann. d. Physik (Wied.), 45, 370-376 (1892).

⁷¹⁰ Coolidge, W. D., Idem, 69, 125-166 (1899).

⁷¹¹ Heerwagen, F., Idem, 49, 272-280 (1893).

Formula (18c) leads to $d\epsilon/dt = -0.383$ for 1 °C at 17 °C and -0.370 at 25 °C, whereas v. Ardenne, Groos, and Otterbein ⁷¹² state that they found -0.36 at 18 °C, and R. King ⁷¹³ states that his observations between 10 and 40 °C (displayed solely as a small-scale graph) satisfy a linear relation, the slope being -0.360 (printed 0.0360).

Values reported by C. B. Thwing ^{713a} and by C. Niven ⁷¹⁴ appear to be untrustworthy. Preliminary measurements at several temperatures below 50 °C have been reported by W. C. Röntgen, ⁷¹⁵ and M. Seeberger ⁷¹⁶ has reported a few measurements, of low accuracy, in the range 15 to 50 °C, $\lambda_0 = 12.6$ to 19.0 cm (see Table 177, note b).

K. Iwanow ⁷¹⁷ has reported that the variation of ϵ with t depends somewhat upon the frequency unless t exceeds 50 °C, and that it is abnormal within an absorption band. He used frequencies between 240 and 500 megacycles/sec, in which region he reported many absorption bands (see Table 173).

The formula here given (18c) and those proposed by the several observers may, for comparison, be thrown into the following forms, in which $\tau \equiv (t-25)/1000$:

$$\epsilon = 78.49 \ | 1 - 4.70_{67} + 10.6_{77}^{2} |$$
 Same as formula (18c)
 $\epsilon = 78.54 \ | 1 - 4.60_{7} + 8.8_{7}^{2} |$ Wyman
 $\epsilon = 78.57 \ | 1 - 4.61_{7} + 15.5_{7}^{2} |$ Drake, Pierce and Dow
 $\epsilon = 78.2e^{-4.7\tau}$ Kockel
 $\epsilon = 78.2_{5}e^{-4.61_{0}\tau}$ Kockel
 $\epsilon = 79.2 \ | 1 - 4.28_{7} + 21.2_{7}^{2} - 410_{7}^{3}$ Linton and Maass
 $\epsilon = 78(1 - 5_{7})$ Linton and Critical Tables
 $n_{t} = n_{25} - 22.0_{7}$; $n = \sqrt{\epsilon}$ Colley

Variation with Pressure.—The dielectric constant of water increases with the pressure. Working at $16.3\,^{\circ}\text{C}$ and $\nu = 60$ megacycles/sec, G. Falckenberg ⁷¹⁸ found that ϵ increased by 0.72 when the pressure was increased from 7 to 200 atm., giving a mean value of $d\epsilon/dp = 0.0037$ cgse per atm.

More recently, S. Kyropoulos,⁷¹⁹ working at 20 °C and $\nu = 0.86$ megacycles/sec, has measured ϵ over the range P = 1 kg*/cm² to P = 3000 kg*/cm². His values are closely given by the empirical formula (19):

$$\epsilon = 80.79 \left[1.0273 + 0.0000372P - 10^{-0.000448(P+3490)} \right]$$
 (19)

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712 v. Aıdenne, Groos, and Otterbein, Physik. Z., 37, 533-544 (1936).
713 King. R., Rev. Sci. Inst., 8, 201-209 (1937).
713a Thwing. C. B., Z. physik. Chem., 14, 286-300 (1894).
714 Niven, C., Proc. Roy. Soc. (London) (A). 85, 139-145 (1911).
715 Röntgen, W. C., Ann. d. Physik (Wied). 52, 593-603 (1894).
710 Seeberger, M., Idem (5), 16, 77-99 (1933).
711 Iwanow, K., Ann. d. Physik (4), 65, 481-506 (1921).
718 Falckenberg, G., Ann. d. Physik (4), 61, 145-166 (1920).
719 Kyropoulos, S., Z. Physik, 40, 507-520 (1926).
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the unit of P being 1 kg*/cm² = 0.9678 atm (see Table 176). It will be noticed that his value (80.79) for 20 °C and 1 atm is about 0.55 per cent greater than the value (80.35) given in Table 174. Using his values of ϵ , the densities (ρ) found by Bridgman (see Table 95), and the molecular weight M=18.00, Kyropoulos computed the value of the molecular polarizability— $M(\epsilon-1)/(\epsilon+2)\rho$ —and of $(\epsilon-1)/\rho$, finding each to vary monotonously throughout the range in P. On going from 1 to 3000 kg*/cm², $M(\epsilon-1)/(\epsilon+2)\rho$ goes from 17.39 to 15.85; and $(\epsilon-1)/\rho$, from 79.9 to 82.6.

An early determination by W. C. Röntgen 715 showed that the excess in the value of ϵ at 500 atm over that at 1 atm is of the order of one per cent.

Table 176.—Variation of the Dielectric Constant of Water with the Pressure

The values (ϵ_k) tabulated by Kyropoulos (see text) are very closely equal to those (ϵ_o) computed by means of the formula $\epsilon_c = 80.79$ (1.0273 + 0.0000372P - E), where $\log_{10}E = -0.000448$ (P + 3490), the unit of P being 1 kg*/cm². For convenience, the values of E and of δ , defined by $\epsilon_o = 80.79$ (1 + δ), are tabulated. Temperature = 20 °C; $\nu = 0.86$ megacycles/sec.

	Unit of $P=1$	$kg*/cm^2 = 0.968$ atm;	of $\epsilon = 1$ cgse unit	
P	e _k	ec	100δ	100 <i>E</i>
1	80.79	80.79	0	2.73
70		80.83	0.05	2.71
206ª		81.83	1.29	2.21
500	83.07	83.18	2.96	1.63
1000	85.20	85.22	5.48	0.97
1500	87.03	87.03	7.73	0.58
2000	88.72	88.73	9.82	0.35
2500	90.34	90.34	11.82	0.21
3000	91.90	91.91	13.77	0.12

^a G. Falckenberg ⁷¹⁸ working at 16.3 °C, $\nu = 60$ megacycles/sec, and p = 7 and 200 atm (7.23 and 206.1 kg*/cm²) found $\epsilon_{200} - \epsilon_7 = 0.72$, as compared with the value 1.00 here given for ϵ_c .

Diclectric Constant of Sca-water.—R. L. Smith-Rose 720 is of the opinion that the dielectric constant of sea-water is about the same as that of distilled water. He attributes the very high apparent values, obtained when audiofrequencies and low radiofrequencies are used, to the effect of polarization films formed at the boundaries between the electrodes and the water.

Dielectric Absorption. (For notation and definitions, see first pages of this Section.)

P. Drude 721 observed that water and certain other substances absorb electric waves much more strongly than can be accounted for by their elec-

⁷²⁰ Smith-Rose, R. L., *Proc. Roy. Soc. (London) (A)*, 143, 135-146 (1933). ⁷²¹ Drude, P., *Ann. d. Physik (Wied.)*, 58, 1-20 (1896); *Z. physik. Chem.*, 23, 267-325 (1897); *Ann. d. Physik (Wied.)*, 64, 131-158 (1898).

trical conductivity. This additional absorption, above that due to the conductivity, is what is here called dielectric absorption. He attributed it to the presence of the hydroxyl group (OH). W. D. Coolidge ⁷¹⁰ referred to the same phenomenon. It arises from a dissipation of energy inherent in the process of polarizing the medium. (See p. 352). For a discussion of the subject on the dipole theory, see P. Debye, ⁷²² and P. Debye and W. Ramm. ⁷²³

A. B. Bryan ⁷²⁴ investigated the phase relations between the current and the emf applied to condensers containing various dielectrics. For water at 23.5 °C and frequencies (ν) of 0.2 to 1.4 megacycles/sec he found that the phase defect was $\phi = 0.8^{\circ} + (2.09/\nu)^{\circ}$, the unit of ν being 1 megacycle/sec. The first term is ascribed to the true dielectric absorption, and the second to the conductivity of the water. It may be easily shown that the part of ϕ arising from the conductivity is $(2/\nu\rho\epsilon)(10^{-6})$ radians = $(114.3/\nu\rho\epsilon)(10^{-6})^{\circ}$, the unit of ν being as before, and the resistivity (ρ) and ϵ being expressed in the same system of units. Equating this to the second term of the observed ϕ , one finds $\rho\epsilon = 55$ microseconds; and if $\epsilon = 81$ cgse unit, $\rho = 0.68 \times 10^{-6}$ cgse unit = 0.6 megohm·cm. The resistivity of best "conductivity" water is nearly 240 megohm·cm, much higher than that used by Bryan.

If the first term of this ϕ arises from dielectric absorption, then its tangent is equal to $\epsilon''/\epsilon' = 2n^2\kappa/\epsilon'$. Hence $2n^2\kappa = \epsilon'$ tan $0.8^\circ = 81(0.0140) = 1.13$, if $\epsilon' = 81$ cgse; whence $\kappa = 0.0070$.

Using better water, J. Granier ⁷²⁵ found $\tan \phi = 0.02$, the portion contributed by the conductivity being 0.006. This gives for $2n^2\kappa$ the same value as was found by Bryan. For the Paris water supply ϕ was 6 times as great.

A. Esau and G. Bäz ⁷²⁶ have studied both the reflectivity and the absorption of water over the range $\lambda_0=2.8$ to 10 cm ($\nu=10,700$ to 3000 times 10^6 cycles/sec), and have presented their results in the form of small graphs. From the heating produced by a field of 28.4×10^6 cycles/sec ($\lambda_0=1055$ cm) C. Schmelzer ⁷²⁷ concluded that the apparent conductivity ($\epsilon''\omega/4\pi$) of the water used ranged from 14 to 20 times 10^{-7} (ohm-cm)⁻¹. Hence ϵ'' ($\equiv 2n^2\kappa$) ranges from 0.09 to 0.13, and κ from 0.0006 to 0.0008, it being assumed that $n^2=\epsilon'=81$. For other determinations see Tables 177 and 178.

The most extended set of values of n and κ now available for water is that by H. W. Knerr. They are given in Table 178 together with certain values derived from them by the compiler. The several values of λ_s have been computed by means of the formulas indicated, numbered as in the text.

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T22 Debye, P., Trans. Faraday Soc., 30, 679-689 (1934); Physik Z., 35, 101-106 (1934).
T23 Debye, P., and Ramm, W., Ann. d. Physik (5), 28, 28-34 (1937).
T24 Bryan, A. B., Phys. Rev. (2), 22, 399-404 (1923).
T25 Granier, J., Bull. Soc. Fr. Élec. (4), 3, 333-482 (1923) = Thesis, Paris.
T26 Esau, A., and Büz, G., Physik Z., 38, 774-775 (1937).
T27 Schmelzer, C., Ann. d. Physik (5), 28, 35-53 (1937).
T28 Knerr, H. W., Phys. Rev. (2), 52, 1057-1067 (1937).
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The discordance between the several sets of values, and their progressive increase with λ_0 , except in the case of (g), indicate that in the case of water the conditions are not so simple as is assumed in deriving the much quoted formulas (8) to (17); that for water we are not justified in assuming that there is only one singularity, and that of the highly dissipative type, in the frequency range below the optical spectrum. This is borne out by the graphs in Figure 6. By formulas (18a) and (18b) each should be

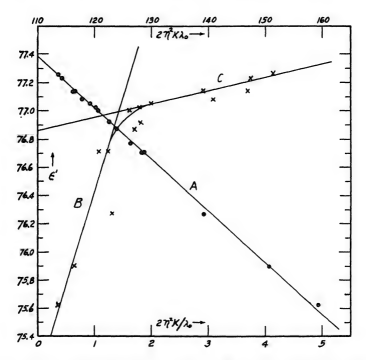


FIGURE 6. Variation of the Dielectric Constant of Water with $2n^2\kappa/\lambda_0$ and with $2n^2\kappa\lambda_0$.

Observations of H. W. Knerr [Phys. Rev. (2), 52, 1054 to 1067 (1937)] at 22 °C. If formulas 18a and 18b were satisfied, the observations would be along two right lines, one cutting the axis of ϵ' at n^2 and the other at ϵ_1 . The observations are linear (dots) in $2n^2\kappa/\lambda_0$, line A, but the intercept is 77.38, whereas ϵ_1 79.61. They are not linear (crosses) in $2n^2\kappa\lambda_0$. See text. The equations of the 3 lines are

(A) ϵ' 77.38 - 0.365 $(2n^2h/\lambda_0)$ (B) ϵ' = 60.50 + 0.1327 $(2n^2h\lambda_0)$ (C) ϵ' - 75.83 + 0.009 $(2n^2k\lambda_0)$.

a right line, and their intercepts on the axis of ϵ' should be ϵ_1 and ϵ_0 . The graph of (18a) is a right line, but its intercept is 77.38, whereas $\epsilon_1 = 79.61$; that of (18b) consists of two right lines, their intercepts being 60.5 and 75.8, whereas $\epsilon_0 = 1.9$.

All of which indicates that a serious doubt attaches to the significance of the several values of λ_8 that have been published, and that the values themselves cannot be satisfactorily compared unless the frequencies and

Table 177.—The Absorption Index of Water

(See also Table 178.)

By definition $I = I_0 e^{-4\pi \kappa n z/\lambda_0}$. The effect of a change in temperature is indicated in footnote b.

	Unit of)	$\omega = 1$ cm; of $\nu = 1$	06 cycles/sec. Te	mp. = 1 °C	
λο	ν	t	K	nĸ	Ref.a
150,000	0.2		0.0070	0.063	Br
21,000	1.4		0.0070	0.063	Br
470	63.8		0.0070	0.063	Gr
242	124	17	0.0032	0.029	Ru
226	133	17	-0.0012	-0.011	Ru
221	136	17	0.0015	0.013	Ru
184	163	17	0.0003	0.003	Ru
160	188	17	0.0020	0.018	Ru
144.5	207.5	17	0.0042	0.038	Ru
144.1	208.0	17	0.0019	0.019	Ru
129	232	17	0.0032	0.029	Ru
106	283	17	0.0093	0.083	Ru
99	303	17	0.0050	0.045	Ru
85	353	17	0.0037	0.033	Ru
75	400	17	0.0114	0.102	Ru
68	441	17	0.0094	0.084	Ru
64	469	17	0.0129	0.116	Ru
57.4	522	17	0.0133	0.119	Ru
57.4	522	17	0.013	0.12	\mathbf{Fr}
52	576	17	0.013	0.12	Fr
23	1300	20.2	0.069	0.62	$\mathbf{Z}\mathbf{a}$
19.0	1580	15	0.079	0.71	Se
14.0	2140	17	0.049	0.45	Se
13.6	2210	16	0.045	0.41	Se
12.6	2380	16	0.058	0.52	Se^b
12.6	2380	15.5	0.059	0.53	Se
8.8	3410	15	0.07	0.60	Ec
5.7	5260	21	0.12	1.08	Ec
4.0	7500	18	0.136	1.18	E 1
3.7	8100	14	0.21	1.72	Ec
2.7	11100	20(?)	0.268	2.26	Te
1.8	16700	20(?)	0.349	2.32	Te
1.75	17100	24	0.35	2.7	Ec
1.5	20000	20(?)	0.276	1.83	Te
1.1	27300	20(?)	0.230	1.44	Te
0.84	35700	20(?)	0.262	1.49	Te
0.42	71400	20(?)	0.240	1.29	Te
" References	s:				
Br Brya Ec Ecke El Elle, Fr Fran Gr Gran Ru Rücl Se Seeb Te Tear Za Zakr	an, A. B. ⁷²⁴ ert, E., I'crh. ph, D., Ann. d. P, kenberger, E., A ier, J. ⁷²⁵ eert, E., Ann. d erger, M. ⁷¹⁶ , J. D., Phys. 1 zewski, K., Bul	. Physik (4), 5: Rev. (2), 21, 61 I. intern. acad.	polon. (A), 1927,	489-503 (1927).	
Seeberger	gives also the	e tollowing to	or $\lambda_0 = 12.6$ cm	:	

15.5

5.9

53 80.6

100κ

 $100n\kappa$

n2

16

5.8

52 80.3 30.0

3.2

28 76.6 50.0

1

9

81

the formulas actually used in computing them are known. Often the formula used is not stated. Some of the published values of λ_{θ} are given in Table 179.

Table 178.—Dielectric Absorption of Water at 22 °C 728 (For definitions and explanation of symbols, see first pages of this Section.)

 $\epsilon = n^2(1 - \kappa^2) - 2in^2\kappa = \epsilon' - i\epsilon''$. In the following formulas (15a), (15b), (15c), and (16), from the text, put $\epsilon_0 = 1.90$ and $\epsilon_1 = 79.61$; in formula (g), similar to (15b), the 77.38 has been obtained from the linear graph in Figure 6. The mean n (n) has been used in all computations.

(15a)
$$\lambda_{s} = \lambda_{0} \left\{ (\epsilon_{1} - \epsilon')/(\epsilon' - \epsilon_{0}) \right\}^{\frac{1}{2}};$$
 (15b) $\lambda_{s} = \lambda_{0} \left(\epsilon_{1} - \epsilon')/2n^{2}\kappa \right)$
(15c) $\lambda_{s} = \lambda_{0} \left\{ 2n^{2}\kappa/(\epsilon' - \epsilon_{0}) \right\};$ (16) $\lambda_{s} = \lambda_{0} \left\{ \frac{A - (A^{2} - 4)^{\frac{1}{2}}}{2} \right\}$
(g) $\lambda_{s} = \lambda_{0} (77.38 - \epsilon')/2n^{2}\kappa;$ $\Lambda \equiv (\epsilon_{1} - \epsilon_{0})/2n^{2}\kappa$

The quantities ν , ϵ' , ϵ'' , ϕ , and λ_8 have been computed by the compiler. Conductivity of the water was 10^{-5} to 10^{-6} (ohm-cm)⁻¹; its contribution to ϵ'' at these frequencies is negligible.

Unit of $\nu = 1000$ megacycles/sec; of $\lambda_0 = 1$ cm; of $\phi = 1^\circ$; of $\lambda_s = 1$ cm $n^2(1-\kappa^2)$ $2n^2\kappa$ 15a 15b g 6.25 4.80 0.153 75.63 17.4 23.7 1.63 1.54 1.11 0.810.3545.62 5.34 0.141 75.90 21.8 16.0 1.58 1.20 0.91 0.362 1.64 8.85 4.63 6.48 0.123 76.27 19.0 14.0 1.70 1.70 1.37 0.378 1.14 3.72 8.05 8.78 0.097 76.71 15.0 1.58 0.360 11.1 1.62 1.61 1.56 3.68 8.16 0.09776.71 15.0 11.1 1.64 1.61 1.58 0.365 1.63 8.77 8.72 3.40 8.80 0.093 76.77 14.4 106 1.69 1.69 1.72 1.73 0.373 3.14 9.55 0.086 76.87 13.3 9.8 1.68 1.69 1.82 1.68 0.366 3.05 9.85 8.80 1.70 1.70 2.97 2.94 10.10 0.082 76.92 12.7 9.3 1.91 2.14 0.366 8.79 10,20 2.76 10.87 0.075 77.00 2.02 2.44 0.357 11.6 8.6 1.66 1.68 2.70 8.83 0.074 8.5 1.69 2.50 11.12 77.02 11.5 1.70 2.07 0.3482.54 11.80 8.85 0.071 77.05 11.0 8.2 1.70 1.72 2.18 2.75 0.3542.24 13.41 77.08 7.7 3.23 0.06810.5 1.851.88 2.450.383 2.07 8.82 14.48 0.06277.14 9.6 7.21.82 1.85 2.62 3.72 0.3611.96 15.29 8.76 0.062 77.14 9.6 7.2 1.93 1.86 2.77 3.93 0 382 1.78 16.83 8.85 77.23 1.63 18.41 0.052 8.0 5.9 1.91 1.95 3.27 5.48 0.346 1.47 20.44 8.78 0.04877.26 7.4 5.5 1.96 2.00 3.62 6.50 0.331 Mean $8.80 = \bar{n}$

Table 179.—Transition Wave-lengths for Water

The transition wave-length (λ_8) is defined as the wave-length in air corresponding to the frequency at which the real part (ϵ') of the dielectric constant is half the sum of the static value (ϵ_1) and the square of the optical index of refraction for infrared waves. There are reasons, for doubting that water fulfills the conditions assumed in deriving the formulas employed

Table 179—(Continued)

in computing λ_s (see text). Knerr's values, computed by the compiler, have been taken from Table 178. The formulas are numbered as in Table 178 and in the text.

All these estimates of λ_s lie within the range 0.36 and 6.5 cm; hence $\lambda_{\tau}(=0.00734\lambda_s$, see p. 356) must lie within the range 0.0026 and 0.048 cm. But the value of τ computed by Debye by means of the oft-quoted formula $\tau = 4\pi\eta r^3/kT$, where η (= 0.01 poise) is the coefficient of viscosity, r (= 2 × 10⁻⁸ cm) is radius of the molecule, k (= 1.327 × 10⁻¹⁶ erg/°K per molecule) is the Boltzmann gas constant, and T (= 293 °K) is the absolute temperature (all referring to water at 20 °C), is τ = 0.27 × 10⁻¹⁰ sec corresponding to λ_{τ} = 0.81 cm—a value that is 17 to 310 times as great as that indicated by the tabulated values.

	Unit of	λ_0 and of $\lambda_0 = 1$ cm.	Temp. = $t ^{\circ}$ C	
λο	t	λε	Eq.	Ref.
5 to 20	22	1.6 to 2.0	16	Knerr
5 to 20	22	1.5 to 2.0	15c	Knerr
5 to 20	22	1.1 to 3.6	15a	Knerr
5 to 20	22	0.8 to 6.5	15b	Knerr
5 to 20	22	0.362	g	Knerr
4	18	1.13	1Ï	Elle
2.8 to 10	19	1.85	15a	Esau and Bäz
1050	20	1.2 to 1.7	10	Schmelzer

[&]quot; References:

Elle, D. See Table 177, references. Esau, A., and Bäz, G., Physik. Z., 38, 774-775 (1937). Knerr, H. W., Phys. Rev. (2), 52, 1054-1067 (1937) $\rightarrow Idem$, 51, 1007 (A₄₆) (1937). (Computation by compiler, see Table 178.) Schmelzer, C., Ann. d. Physik (5), 28, 35-53 (1937).

50. CONDUCTION OF ELECTRICITY BY WATER

The National Research Council has awarded to G. A. Hulett a grant for a study of the electrical conductivity of pure water.⁷²⁹

The observation of O. Risse, 730 that when water is exposed to x-rays its conductivity is increased and its pH is decreased, is probably to be explained by the heating of the water by thermal radiation from the anticathode. H. Fricke and E. R. Brownscombe, 732 using methods more sensitive than those employed by Risse, were unable to detect any formation of either H_2O_2 or O_2 when water is exposed to x-rays, even though as much as 150 kiloroentgens were used.

All the following data regarding the electrical conduction of water, with the exception of those for natural waters (p. 380 and Table 184) and those published since 1929, have been obtained from the *International Critical Tables*, either directly or by means of formulas there given.

⁷²⁰ Hulett, G. A., Science, 77, 215 (1933).

⁷³⁰ Risse, O., Z. physik. Chem. (A), 140, 133-157 (1929).

⁷⁸¹ Schnurmann, R., Idem, 150, 110-114 (1930).

⁷⁸² Fricke, H., and Brownscombe, E. R., Phys. Rev. (2), 44, 240 (1933).

For the purposes of this compilation, the data have been grouped under three heads: (1) Conductivity of water, (2) Equivalent conductivity of the ions, (3) Electrolytic ionization.

Conductivity of Water.

The conductivity (κ) of a substance is the reciprocal of its volume resistivity, which is the longitudinal resistance per unit of length of a uniform cylinder of the substance of unit cross-sectional area. The dimensions of κ are given by the formula

$$\frac{length}{(area \times resistance)} = (resistance \times length)^{-1}$$

This quantity κ is also called both the specific conductance and the volume conductivity.

The best water obtained by Kohlrausch and Hevdweiller is reported to have had at 10 °C the conductivity $\kappa = 4.2 \times 10^{-8}$ (ohm·cm)^{-1,733} This is about 10 per cent greater than that computed from the equivalent conductivities of the ions and the electrolytic ionization of water (see Table 180).

Table 180.—Electrical Conductivity of Pure Water (Computed)

The electrical conductivity (κ) of pure water is related to the equivalent conductivities of its ions (Λ_{II} and Λ_{OII}), its density (ρ), and its ionization product (K) as indicated by the equation $\kappa = (\Lambda_{\rm H} + \Lambda_{\rm OH})_{\rho N}/\overline{K}/1000$. The following three sets of values of κ have each been computed from the values of $\Lambda_{\rm H}$ and $\Lambda_{\rm OH}$ as given in Table 181, the values of ρ as given in Table 93, and the values of K as defined (see N. Bjerrum 733) in one of the following ways: (1) by the "best" values of p_w (see ICT of Table 182); (2) by the formula of A. Heydweiller, $^{738} \log_{10} K^{-1} = 6099.6/(273 + t) +$ $24.25 \log_{10}(273 + t) - 66.4678$; or (3) by the formula of G. N. Lewis and M. Randall, $^{739} \log_{10} K^{-1} = 6384.7/(273.1 + t) + 26.676 \log_{10}(273.1 + t) -$ (For definition of Λ and of K, see following paragraphs.)

		Unit of $\kappa = 1$	(onm·cm)-1.	Temp. $= t$ °C		
$K \rightarrow$	(1) ·		(2)		(3)	
ı	log ₁₀ κ	10 ⁸ κ	Log ₁₀ ^K	$10^{8}\kappa$	Logior	10 ⁸ κ
0°	8.07_{5}	1.19	$\bar{8}.064$	1.16	$\bar{8}.067$	1.17
10	8.36_{3}	2.3_{1}	$\overline{8}.357$	2.28	8.360	2.29
15	8.49_{7}	3.14	8.491	3.10	8.493	3.11
18	8.57_{4}	3.75	8.568	3.70	8.569	3.71
25	8.74_1	5.5_{1}	8.736	5.44	8.736	5.44
35	$\overline{8}.96_1$	9.14	8.954	9.00	8.952	8.96
40	$\bar{7}.06_{0}$	11.5	$\bar{7}.051$	11.25	$\overline{7}.048$	11.17
50	7.234	17.1	$\bar{7}.232$	17.06	7.226	16.83

⁷³³ Bjerrum, N., Int. Crit. Tables, 6, 152 (1929), from Kohltausch, F., and Heydweiller, A., Ann. d. Physik (Wied), 53, 209-235 (1894); Z. physik. Chem, 14, 317-330 (1894); also, Partington, J. R., Int. Crit. Tables, 6, 142 (1929).

⁷³⁴ Kling, A., and Lassieur, A., Ann. de Chim. (10), 15, 201-227 (1931); Compt. rend, 201, 203-204 (1935); Jour. de Phys. (7), 7, C.P. 21 (1936) ← Doc. Sci., 4, 225-229 (1935). 785 Gostkowski, K., Z. physik. Chem. (A), 170, 149-152 (1934); Acta Phys. Polon., 3, 75-80

⁷³⁶ Thiessen, P. A., and Hermann, K., Z. Elektroch., 43, 66-69 (1937).

In recent years A. Kling and A. Lassieur ⁷³⁴ have persistently maintained that the exact conditions necessary for obtaining water of minimum conductivity are not known, and have presented experimental evidence which they think justify that conclusion. But the lowest conductivity that they reported (2×10^{-6}) was much greater than that found by Kohlrausch and Heydweiller.

K. Gostkowski ⁷³⁵ has described a paraffin-lined still from which, by low temperature distillation, he has obtained water for which $10^8\kappa = 7 \, (\text{ohm} \cdot \text{cm})^{-1}$ at $0 \, ^{\circ}\text{C}$. And P. A. Thiessen and K. Hermann ⁷³⁶ have described a procedure by which they obtained 400 cm³/hr of water having $10^8\kappa = 6.5$ to 8 $(\text{ohm} \cdot \text{cm})^{-1}$ at 25 $^{\circ}\text{C}$, and have stated that by the use of an additional stage in the distillation they got $10^8\kappa = 5.95 \, (\text{ohm} \cdot \text{cm})^{-1}$ at 25 $^{\circ}\text{C}$; they give the "theoretical" value as 5.52.

A. Deubner ^{7:37} has reported that the conductivity of water in contact with air may decrease from $10^6 \kappa = 6$ or 8 (ohm·cm)⁻¹ as the water left the still, to 1.5 (ohm·cm)⁻¹ after it had been in contact with air for about a day. The explanation is not known, but he considered several possibilities.

Equivalent Conductivity.

The equivalent conductivity (Λ) of an electrolytic solution is defined by the relation $\Lambda = \kappa/c$, where c, called the equivalent concentration, is the number of equivalents (more explicitly, electrolytic equivalents) of the solute per unit volume of the solution. The equivalent conductivity is the axial conductance of a cylindrical volume of the solution, of unit length and of such a cross-sectional area that it contains one equivalent of the solute. Its dimensions are given by the formula:

$$\frac{\text{volume per equivalent}}{\text{resistance} \times \text{length}} = \frac{\text{area per equivalent}}{\text{resistance}}$$

The electrolytic, or electrochemical, equivalent of an ion is defined as the total mass of such ions needed to carry a combined resultant charge equal to a unit quantity of electricity. It may be expressed in terms of any one of various units, such as a gram per coulomb, or a gram per faraday. In the following, as in most physicochemical work, the second of these units will be employed, a faraday being understood to denote the resultant charge carried by one gram-formula weight of a univalent ion, which charge is not far from 96,500 coulombs.⁷⁴⁰ When expressed in that unit, the electrolytic equivalent of an ion is equal to the formula-weight of the ion divided by its valence. Likewise, the electrolytic equivalent of a solute will be understood to mean the formula-weight of the solute divided by the combined

⁷³⁷ Deubner, A., Ann. d. Physik (4), 84, 429-456 (1927).

⁷⁸⁸ Heydweiller, A., Ann. d. Physik (4), 28, 503-512 (1909).

⁷³⁰ Lewis, G. N., and Randall, M., "Thermodynamics and the free energy of chemical substances," New York, 1923.
740 Int. Crit. Tables, 1, 17 (1926).

valences of the similarly charged ions to which each formula-molecule may give rise.

If Λ is the equivalent conductivity of a solution in which the total equivalent concentration of the similarly charged ions is c_0 , and if c_1 , c_2 , c_3 , are the equivalent concentrations of the several species of ions to which the solute actually gives rise, then it is possible to assign to each ion a quantity $(\Lambda_1, \Lambda_2, \Lambda_3, \ldots)$, called its equivalent conductivity, which depends only upon the temperature, upon the natures of the ion and of the solvent, and upon the value of c_0 ; which satisfies the formula $\Lambda c_0 = \Lambda_1 c_1 + \Lambda_2 c_2 + \Lambda_1 c_1 + \Lambda_2 c_2 + \Lambda_2 c_2 + \Lambda_3 c_2 + \Lambda_3 c_3 + \Lambda_4 c_4 + \Lambda_5 c_5 + \Lambda_5$ $\Lambda_{3}c_{3} + \dots$; and which approaches a definite limit as c_{0} is reduced indefinitely, that limit being characteristic of the ion and independent of the natures and numbers of the other species of ions present. If the solute, which may be a mixture of substances, is completely dissociated, then $c_0 = c$, the equivalent concentration of the solute; in other cases, $c_0 < c$. If there is only a single solute and if it is completely dissociated, giving rise to only two species of ions, then $c_1 = c_2 = c$ and $\Lambda = \Lambda_1 + \Lambda_2$. This is the case for pure water, in which the molecules (taken as H₂O) that are ionized are regarded as those of a solute, the others as those of the solvent. case the ions are H+ and OH-, and the concentration is always very low (see next paragraph), so low that Λ_1 and Λ_2 have practically their limiting, constant values corresponding to zero concentration.

The relation of the structure of the molecule to the conductivity of water has been considered by many, some of the more recent being J. D. Bernal and R. H. Fowler,⁷⁴¹ J. D. Bernal,⁷⁴² John Rehner, Jr., ⁷⁴³ and G. Wannier.⁷⁴⁴

Table 181.—Equivalent Conductivities of the Ions of Water. 745

Unit of $\Lambda = 1$ cm²·equivalent⁻¹ ohm⁻¹; temp = t°C. Concentration indefinitely low $t \qquad \qquad \Lambda_H \qquad \qquad \Lambda_{OH} \qquad \qquad \Lambda_H + \Lambda_{OH}$

4	Λн	Аон	Лн +Лон
0	229.0	118	347.0
10	275.6	149	424.6
15	300.4	164.5	464.9
18	315.2	1740	489.2
25	350.0	196	546.0
35	399.6	228	627.6
40	421.4	244	665.4
50	464.3	276	740.3

° In the introduction to the section on Electrical Conductivity of Aqueous Solutions, 760 this value is given as 173.8, making $\Lambda_{\rm H}+\Lambda_{\rm OH}=489.0$. At the same place and for 18 °C the following values are given also:

$$\frac{1}{\Lambda_{\rm H}} \left(\frac{d\Lambda_{\rm H}}{dt} \right) = 0.01573, \ \frac{1}{\Lambda_{\rm OH}} \left(\frac{d\Lambda_{\rm OH}}{dt} \right) = 0.018 \ \text{per} \ ^{\circ}\text{C}.$$

⁷⁴¹ Bernal, J. D., and Fowler, R. H., J. Chem'l Phys., 1, 515-548 (1933).

⁷⁴² Bernal, J. D., Trans. Faraday Soc., 30, 787 (1934).

⁷⁴⁸ Rehner, John, Jr., Rev. Sci. Inst. (N. S.), 5, 2-3 (1934).

⁷⁴⁴ Wannier, G., Ann. d. Physik (5), 24, 545-568, 569-590 (1935) = Diss., Basel.

⁷⁴⁵ Kendall, J., Int. Crit. Tables, 6, 259-304 (259) (1929).

⁷⁴⁶ Anon, Int. Crit. Tables, 6, 230, Table 3 (1929).

Electrolytic Ionization of Water.—(For photochemical dissociation and the energy involved in ionic dissociation, see Section 8.)

Let the symbol [H+] denote the number of moles of H+ ions per 1000 units of mass of water*; that is, per 1000/18.0154 moles of H₂O; and similarly for [OH-]. If water dissociates solely in the manner H₂O ≠ $H^+ + OH^-$, then for pure water, $[H^+] = [OH^-]$. If the addition of a solute adds H+ ions to the solution, some of these will combine with OHions to form neutral molecules. This will continue until the product K =[H⁺]·[OH⁻] is the same as it would be for pure water, but now, [H⁺] does not equal [OH-]. Similarly if the solute adds OH- ions. (A. Kling and A. Lassieur 747 have suggested that the dissociation of water may not be restricted to the type just mentioned.)

The quantity K is called the ionization product, and $p_w = \log_{10} K^{-1}$ is called the ionization exponent of water.

From these definitions and those in the earlier portions of this section, it follows at once that for pure water $1000c/\rho = [H^+] = [OH^-] = \sqrt{K}$,

and $\frac{1}{1000c} = \frac{1}{\rho\sqrt{K}}$ where ρ = density. If the unit of mass = 1 gram and

of volume = 1 milliliter, then 1000c is the number of gram-equivalents of either ion per liter of pure water.

The symbol pH is commonly used to denote the common logarithm (base 10) of the reciprocal of the number of gram-equivalents of H+ per liter of solution (i.e., of the reciprocal of the "hydrogen-ion concentration"). Hence, for pure water pH = $\log_{10}(\rho\sqrt{K})^{-1} = 1/2p_w - \log_{10}\rho$ if the units are those just stated.

E. Truog 748 has stated that, for water at 25 °C in equilibrium with air of average CO₂ content, pH = 5.7 to 5.8, and not 7, as frequently assumed. This value is affected but little by changes in temperature, the effect of changes in the solubility of CO₂ being largely offset by changes in the ionization. But if the air has been carefully freed of CO2 and of NH3, the pH of water at 25 °C will be about 7. The time required for exposed water to come into equilibrium with the CO₂ in the air is very brief (see Section 86).

Similarly S. B. Ellis and S. J. Kiehl 749 found for the purest water pH = 7.01 at 27.5 °C; and that value has been confirmed by J. A. Cranston and H. F. Brown. 750

^{*}Although it is customary and convenient to define the quantity [H⁺] either in this way or in terms of a liter instead of a kilogram, such a definition introduces a purely arbitrary numerical factor. It would be more consistent with sound scientific custom to define [H⁺] as the ratio of the number of moles of H⁺ ions to the total number of moles (i.e., of formula-weights) of H₂O, and similarly in other cases.

⁷⁴⁷ Kling, A., and Lassieur, A., Compt. rend., 181, 1062-1064 (1925); Ann. de Chim. (10), 15, 201-227 (1931).

⁷⁴⁸ Truog, E., Science (N. S.), 74, 633-634 (1931).

⁷⁴⁰ Ellis, S. B., and Kiehl, S. J., J. Am. Chem. Soc., 57, 2145-2149 (1935).

⁷⁵⁰ Cranston, J. A., and Brown, H. F., Trans. Faraday Soc., 33, 1455-1458 (1937).

In striking contrast with other observers, A. Kling and A. Lassieur 751 have reported the very low value pH = 5.8, and maintain it in the face of criticism by R. Cliquet-Pleyel 752 although the conductivity of their water was very high. 758

Table 182.—Ionization Exponent and Product for Water

(For comparison of experimental values, see Table 183.)

In computing the H and the LR values, the logarithms were carried one place farther than they are given in this table. The tabulated values of K correspond to the more exact logarithms, whence such apparent discrepancies as occur between K and p_w at 25 °C. For the pH value of water in contact with air, see text.

K	=[H ⁺]·[OH ⁻]	. Unit of [H+]	and of [OH-]=	1 g-mole of ion	per 1000 g wate	er
Sourcea→	ICT	$H = p_{so} = \log_{10} K^{-1}$	LR	ICT	H 1014K	LR
o	14.93	14.952	14.946	0.117	0.112	0.113
5	14.72	14.741	14.736	0.190	0.182	0.184
10	14.53	14.541	14.537	0.295	0.287	0.291
15	14.34	14.352	14.348	0.457	0.445	0.448
18	14.23	14.242	14.240	0.589	0.572	0.576^{b}
20	14.16	14.171	14.170	0.692	0.674	0.677
25	13.99	14.000	14.000	1.02	0.999	1.000^{b}
30	13.83	13.838	13.840	1.48	1.45	1.45
35	13.67	13.684	13.687	2.14	2.07	2.06
37	13.61	13.624	13.628	2.46	2.38	2.35
40	13.52	13.537	13.543	3.02	2.91	2.87
45	13.39	13.397	13.406	4.07	4.01	3.93
50	13.26	13.265	12.276	5.50	5.44	5.30
60	13.03	13.019	13.036	9.33	9.58	9.21
70	12.82	12.796	12.820	15.1	16.0	15.2
80	12.63	12.595	12.626	23.4	25.4	23.7
90	12.45	12.413	12.451	35.5	38.6	35.4
100	12.29	12.249	12.295	51.3	56.4	50.8
150	11.63	11.641	11.729	234	229	186
200	11.26	11.293	11.428	550	509	373
250	11.17	11.119	11.302	676	761	499
300	11.40	11.062	11.295	398	866	507
306	11.46	11.062	11.300	347	867	501
		Harne	ed and Geary	(HG)a		
Sourcea→	нн,нс	HM	HD	HG	Mean	Mean
			1014K			logicK-1
0	(0.115)	0.1134	0.1132	0.1125	0.1133	14.9458
5	(0.186)	0.1850	0.1842	0.1834	0.1846	14.7333
10	(0.293)	0.2919	0.2921	0.2890	0.2920	14.5346
15	(0.452)	0.4505	0.4504	0.4500	0.4503	14.3465
20	0.681	0.6806	0.6806	0.6815	0.6809	14.1669
25	1.008	1.007	1.007	1.009	1.008	13.9965
30	1.471	1.470	1.467	1.466	1.468	13.8333
35	2.088	2.091	2.088	2.090	2.089	13.6801
40	2.916	2.914	(2.891)	2.920	2.917	13.5351
45	4.016	4.017		4.023	4.018	13.3960
EO	E 176	E 102		FACE	E 474	42000

na Kling, A., and Lassieur, A., Ann. de Chim. (10), 15, 201-227 (1931); Compt. rend., 201, 203-204 (1935).

Table 182—(Continued)

" Sources:

Bjerrum, N.,783 those designated as "best values"; they are based upon the experimental values listed in Table 183. Values for temperatures below 100 °C rest upon no determination of potential or of hydrolysis based upon dissociation as inferred from the conductivity; at higher temperatures the greatest weight is given to the determination of potential. ICT

A. Heydweiller's formula ⁷³⁸ (see also Bjerrum, N.⁷³⁸), $\log_{10}K^{-1} = \frac{6099.6}{273 + t} + 24.25$ н $\log_{10}(273+t)-66.4678.$

HC

HDHG

HH

log₁₀(2/3 + 1) = 00.40/8. Harned, H. S., and Copson, H. R., J. Am. Chem. Soc., 55, 2206-2215 (1933). Harned, H. S., and Donelson, J. G., Idem, 59, 1280-1284 (1937). Harned, H. S., and Geary, (*. G., Idem, 59, 2032-2035 (1937). Harned, H. S., and Hamer, W. J., Idem, 55, 2194-2206, 4496-4507 (1933). Harned, H. S., and Mannweiler, G. E., Idem, 57, 1873-1876 (1935). Formula given by G. N. Lewis and M. Randall. (See Bjerrum, N. 783), log₁₀K⁻¹ = HM LR

 $\frac{6384.7}{273.1+t} + 26.676 \log_{10}(273.1+t) - 73.424.$

 $^{\circ}$ In his compilation, 754 M. Randall gives for the "equilibrium constant (activities)" $\times 10^{14}$ the values 0.114 at 0 °C, 0.58 at 18 °C, and 1.005 at 25 °C, based on the data of G. N. Lewis and M. Randall, 766 and of R. Lorenz and A. Böhi. 766

Table 183.—Ionization Exponent for Water: Comparison of Values

For final value derived by Bjerrum, and for values computed by means of formulas, see Table 182.

All data in this table have been taken from the compilation of N. Bjerrum.⁷³³ Recently, E. J. Roberts (Ro)⁶ has reported for 25 °C, $10^{14} K = 0.988 \pm 0.004$; i.e., $\log_{10} K^{-1} = 14.005 \pm 0.002$.

Meth	ode Cond		Hwd	rolysis -				Dote	ntial		
Prope		- Ioniza		Act	ivity —		Ion-cone	entration	TILIAI —	Acti	vity —
Ref.		Lun	Kan	Lun	Kan	LB	PT	Mic	Sor'	LRd	Bu*
ı	1100		4 - 0 -		44.00	$p_w = \log$	101 -			44.045	44.006
.0	14.93		15.05		14.99	14.87				14.945	14.926
10	14.52	14.51		14.47							
15		14.34		14.30							
18	14.22		14.34		14.27	14.15		14.13	14.14	14.239	14.222
25	13.98	13.98	14.09	13.94	14.03	13.92	13.91	13.89		13.998	13.980
30						13.76		13.72			
37								13.50		13.626	13.590
40		13.53		13.49		13.41		13.42			
50	13.25	13.29		13.25		13.06				13.273	
60						12.90					
70						12.67					
80						12.46					
90		(NK)	c			12.37					
100		12.28				12.07				12.29	
156		11.57	(Sos)	:						12.27	
		11.57									
218			11.19								
306			11.46								

[&]quot;Methods: Cond = computed from the conductance of the purest water. The values are said to have been computed by means of the Hcydweiller formula (H of

⁷⁰² Cliquet-Pleyel, R., Jour. de Phys. (7), 7, C.P. 21 (1936) \leftarrow Doc. Sci., 4, 104-113 (1935); Chem. Abst., 31, 7727 (1937) \leftarrow Doc. Sci., 5, 65-70 (1936).

⁷⁰³ Kling, A., and Lassieur, A., Jour. de Phys. (7), 7, C.P. 21 (1936) ← Doc. Sci., 4, 225-229 (1935); Lassieur, A., Chem. Abs., 31, 7727 (1937) ← Doc. Sci., 5, 11-15 (1936).

704 Randall, M., Int. Crit. Tables, 7, 224-313, 347-353 (232) (1930).

⁷⁶⁵ Lewis, G. N., and Randall, M., J. Am. Chem. Soc., 36, 1969-1993 (1914).

⁷⁵⁶ Lorenz, R., and Böhi, A., Z. physik. Chem., 66, 733-751 (1909); cf. G. N. Lewis and M. Randall. 759

Table 183—(Continued)

Table 182), but they differ appreciably from those so computed by the present compiler (see Table 182).

Hydrolysis = computed from the hydrolysis of weak acids and of weak bases. Potential = computed from the potentials of cells having a hydrogen electrode in either an acid or an alkaline solution.

Property utilized: Both ionization and ion-concentration were determined from the observed conductance. Activity = activity-coefficient (f) of the ions; in the method of hydrolysis, it was computed by means of the formula: $\log_{10} f^{-1} = 0.3 \sqrt{c_{i}}$; $c_i = \text{concentration of either species of ion (unit = 1 g-mole per kg of water)}$; in the method of potential, it was calculated by thermodynamic methods.

References:

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9, No. 1, pp. 208 (1929).
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Lewis, G. N., and Randall, M. Tao
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Michaelis, L., "Die Wasserstoffionenkonzentration," Berlin, J. Springer, 1914.
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Poma, G., and Tanzi, B., Z. physik. Chem. 79, 55-62 (1912).
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Mic NK

PT Sor

"Observations were made at 25 °C, and values of K at other temperatures were computed from that at 25 °C by means of the LR formula of Table 182, which assumes that the heat of neutralization is (29210-53T) cal₂₀ per equivalent.

The Debye-Hückel square-root formula was assumed as the limiting law at infinite dilution. $\Lambda = f_{\lambda}\Lambda_{0}$, $f_{\lambda} = 1 - A\sqrt{c_{1}}$ where A depends upon the solvent, the solute, and the temperature, and $c_4 =$ concentration of either species of ion; the unit of c_i must accord with those used in A as $A\sqrt{c_i}$ is dimensionless.

Conductivity of Rain-water.

For freshly fallen rain-water at 17.6 °C, H. Schmidt ⁷⁵⁸ found $10^6 \kappa = 128$ (ohm·cm)⁻¹. For purest water at 17.6 °C, $10^6 \kappa = 0.036$ (ohm·cm)⁻¹, Table 180.

Conductivity of Sea-water.

The composition of sea-water varies from place to place, and from time to time, depending upon the evaporation and the inflow of fresher water from streams, ice-bergs, and precipitation; its electrical conductivity likewise varies. Sea-water averages about 35 g of salts per kg. Data for variations in the salinity (S), composition of the salt, and temperatures of the oceans are given elsewhere (see p. 654). In the following, the unit of S is 1 g salt per kg of sea-water.

E. Ruppin 759 has found that the electrical conductivities at 0, 15, and 25 °C, respectively, may be computed by means of the expressions: $10^6 \kappa_0 =$ $978S - 5.96S^2 + 0.0547S^3$; $10^6 \kappa_{15} = 1465S - 9.78S^2 + 0.0876S^3$; $10^6 \kappa_{25} =$ $1823S - 12.76S^2 + 0.1177S^3$; and O. Krümmel 700 has stated that Knudsen

⁷⁵⁷ Debye, P., and Hückel, E., Physik. Z., 24, 305-325 (1923).

⁷⁵⁸ Schmidt, H., Jahrb. d. drahtlos. Teleg., 4, 636-638 (1911).

⁷⁵⁰ Ruppin, E., Wiss. Meeresunters. (N. F.), 9, (Abt. Kiel), 178-183 (1906) $\rightarrow Z$. anorg. Chem., 49, 190-194 (1908).

⁷⁰⁰ Krümmel, O., "Handb. d. Ozeanog.," Vol. 1, 1907.

has found that, whatever the value of S, $\log_{10\kappa_t} = \log_{10\kappa_{15}} + \alpha(t-15)$, where $\alpha = 0.01135$ when t = 0 °C, and $\alpha = 0.00928$ when t = 25 °C.

The conductivity is the same for all frequencies from zero to 100 kilocycles/sec,⁷⁶¹ but R. L. Smith-Rosc ⁷²⁰ has reported that the conductivity of a sample taken from the English Channel increased, at 20 °C, from $0.043 \text{ (ohm \cdot cm)}^{-1}$ at 0.5 kc/sec to $0.060 \text{ (ohm \cdot cm)}^{-1}$ at 10.000 kc/sec. He found a mean temperature coefficient of k to be 2.7 per cent per 1 °C, between 0 and 40 °C; Rivers-Moore 761 derived three per cent from observations in the range 12.0 to 18.3 °C; and H. Schmidt 758 derived 1.5 from observations in the range 18 to 21.4 °C. All of these are markedly greater than Knudsen's values, and rest on less extensive data.

Table 184.—Electrical Conductivity of Sea-water 760

Accepting Ruppin's formulas (see text) connecting k and the salinity (S) and Knudsen's values for the temperature coefficient (α) of κ , and assuming that the variation in a is linear in the temperature, Krümmel has computed the following values.

Unit of S=1 g salt per kg sea-water; of $\kappa=1$ (ohm-cm)⁻¹. Temp. = 1°C

\rightarrow	5	10	15	20 10	25	30	35	40
)	48	92	135	176	216	254	293	331
5	55	107	156	203	248	292	335	378
)	63	122	178	231	283	332ª	382	430
5	71	138	201	261	319	375	431	486
)	79	154	225	292	357	420b	482	543
5	88	171	249	323	394	464	532°	601
Ď	97	187	273	354	433	510	585	660

^a For sea-water from off the coast near Hastings, England, S not stated, Balth.v.d.Pol ⁷⁰¹ found $10^4\kappa = 377$ at 12.5 °C.

^b For surface water from the North Sea, S not stated, H. Schmidt ⁷⁰⁸ found $10^4\kappa = 397$ at 20 °C.

^c E. G. Hill ⁷⁰² measured κ at 25 °C for sea-water of various concentrations lying between C1 = 20.439 (S = 37) and (C1 = 21.533 (S = 39), obtaining values about 2.5 per cent smaller than those given by Ruppin's formula. But T. Shedlovsky ⁷⁰⁸ found for a sample from Tortugas, 25 °C, Cl = 19.92 g/kg (S = 36.10), $\kappa = 0.05419$, only 0.8 per cent smaller than the tabular values indicate.

51. KERR ELECTRO-OPTIC EFFECT FOR WATER

When water is subjected to a uniform electric field it becomes slightly birefringent, behaving like a positive uniaxial crystal with its axis in the direction of the field. If n_0 and n_0 denote, respectively, the ordinary and the extraordinary index of refraction for light of wave-length λ in vacuo, then $(n_0 - n_0)/\lambda E^2 = C$ is independent of the field (E) and is known as the Kerr electro-optic constant, or the coefficient of electric birefringence. The value of C depends upon λ and the temperature as well as upon the

 ⁷⁶¹ v. d. Pol, Balth., Jr., Phil. Mag. (6), 36, 88-94 (1918); Rivers-Moore, H. R., Electrician (London), 82, 174-176 (1919).
 762 Hill, E. G., Proc. Roy. Soc. Edinburgh, 27, 233-243 (1907).

⁷⁰³ Shedlovsky, T., private communication from Dr. L. R. Blinks through Mr. R. S. Ould.

nature of the substance. T. H. Havelock ⁷⁶⁴ has developed a theory that seems to accord well with the observed facts. On this theory $C\lambda n/(n^2-1)^2$ is a constant for a given temperature, n being the ordinary index of refraction for light of wave-length λ . This theory, as well as that of Cotton and Mouton, requires that $(n_e-n)/(n_0-n)=-2$. The experimental determination of this ratio of the absolute retardations is rendered extremely difficult by the presence of electrostriction and of thermal effects. In those cases in which the effect of these has been eliminated, the ratio has been found to be -2.765 Apparently it has not been determined for water.

M. Pauthenier ⁷⁶⁶ reports that at 17 °C and for the D-line the value of C for water is 1.23 times its value for CS_2 . This is the value accepted by H. Mouton ⁷⁶⁷ as the best available at the time his compilation was prepared. Accepting, with Pauthenier, L. Chaumont's data for CS_2 , ⁷⁶⁸ Havelock's formula for the variation of C with λ and n, and F. Martens' formula for the refraction of CS_2 , ⁷⁶⁹ increasing that n by 0.00090 in order to reduce it from 18° to 17 °C, we have as follows for CS_2 at 17 °C: For $\lambda = 0.54607 \,\mu$, $10^7C = 3.6315$, n = 1.63811, $10^{11}C\lambda n/(n^2 - 1)^2 = 1.1463$; for $\lambda = 0.57801 \,\mu$, $10^7C = 3.3580$, n = 1.63171, $10^{11}C\lambda n/(n^2 - 1)^2 = 1.1459$. Using the mean value $10^{11}C\lambda n/(n^2 - 1)^2 = 1.1461$; we find for $\lambda = 0.58931 \,\mu$ and $n_{17} = 1.62974$, $10^7C = 3.2728$ for CS_2 , the D-lines, and 17 °C. This essentially agrees with the value that may be derived from the data given in Mouton's compilation. ⁷⁶⁷

Whence for water at 17 °C and the D-lines ($\lambda = 5893A$)

$$C = 4.03 \, 10^{-7} \text{cgse}$$

= $0.0363 \, \text{cm/volt}^2$

Taking n=1.33324, this gives for the Havelock constant for water at 17 °C the value $C\lambda n/(n^2-1)^2=5.24$ $10^{-11}{\rm cgse}=4.72$ $10^{-6}{\rm cm}^2/{\rm volt}^2$. In his review of the molecular field problem, F. G. Keyes ⁷⁷⁰ accepts the value (reference not given) $C\lambda/n=14.4$ $10^{-12}{\rm cgse}$ for water at 20 °C. That leads to C=3.25 $10^{-7}{\rm cgse}$ for the D-lines and water at 20 °C, which can be reconciled with the preceding value for 17 °C only by assuming the very great temperature coefficient of 8 per cent per 1 °C. That for CS₂ is only 0.5 per cent per 1 °C; using that with the 17 °C value for C leads to 17.6 $10^{-12}{\rm cgse}$ for $C\lambda/n$ for water at 20 °C, which is the value given by P. Debye ⁷⁷¹ on the basis of Pauthenier's observations. (For the quantity $C\lambda/n$, 1 cgse = 1 ϵ ·cm³/erg, ϵ = the unit of dielectric constant; for C, 1 cgse = 1 ϵ ·cm²/erg.)

If the electric field is oscillatory and its frequency is near that of a

⁷⁶⁴ Havelock, T. H. Proc. Roy. Soc. (London) (A), 77, 170-182 (1905); 80, 28-44 (1907); Phys Rev., 28, 136-139 (1909).

⁷⁶⁶ Pauthenier, M., Ann. de Phys. (9), 14, 239-306 (1920); Jour. de Phys. (6), 2, 183-196 (1921).

⁷⁰⁰ Pauthenier, M., Jour. de Phys. (6), 2, 384-389 (1921).
707 Mouton, II., Int. Crit. Tables, 7, 109-113 (110) (1930).

⁷⁰⁸ Chaumont, L., Ann. de Phys. (9), 4, 61-100, 101-206 (1915); 5, 17-78 (1916).

⁷⁰⁰ Martens, F. F., Ann. d. Physik (4), 6, 603-640 (632) (1901).

⁷⁷⁰ Keyes, F. G., Chem'l Rev., 6, 175-216 (1929).

⁷⁷¹ Debye, P., Handb. d. Radiol. (Marx), 6, 597-786 (770) (1925).

characteristic system in the substance, the Kerr constant may differ markedly from its value for static fields. Effects, at first thought to be of this kind, were reported for water by A. Bramley,⁷⁷² but were later shown by him to have another origin.⁷⁷⁸

52. ELECTRICAL DISCHARGE IN WATER

High-potential discharges between electrodes immersed in water are of various types—arc, brush, corona, spark—depending upon the nature of the circuit, and are accompanied by mechanical disturbances of the liquid. Such disturbances may exist—owing to electrostatic forces between the electrode and the electrically charged water—even when the discharge is feeble. M. Katalinic ⁷⁷⁴ has described the production of waves and of sprays when a high-voltage a.c. potential is applied to a wire electrode dipping in, or lying just below, the surface. He states that at 0.0004 cm from the electrode the potential gradient may be over 2 megavolts/cm when the applied voltage is 1200 volts.

Arc.

The spectrum of under-water arcs between carbon electrodes has been studied by H. Konen 775; and H. D. Carter and A. N. Campbell 776 have studied the electrical products formed, using electrodes of various materials. The latter found that the rate of evolution of gas increases with the temperature of the water, and is independent of the pressure, at least to 23 atm, and that the temperature of the arc depends upon the nature of the electrodes and upon the temperature of the water, increasing about 1500 °C when the temperature of the water is increased from 5 °C to 100 °C. They give a bibliography of 46 entries. J. W. Shipley 777 has studied the arcing that occurs under certain conditions in the a.c. electrolysis of water, using solutions of NaOH.

Brush.

The color and spectrum of the brush discharge in water, and their variations with the conditions, have been studied by H. Smith.⁷⁷⁸

Corona.

Two types of impulse corona in water have been described by Y. Toriyama and U. Shinohara.⁷⁷⁹ Using point-to-plate electrodes and impulses of 10 to 100 microseconds duration, they obtained a pink discharge having a line spectrum. With a higher crest voltage and a duration of only 0.1

⁷⁷² Bramley, A., J. Franklin Inst., 206, 151-157 (1928); Phys. Rev. (2), 33, 640 (1929).

⁷⁷⁸ Bramley, A., J. Opt. Soc. Amer., 21, 148 (1931).

⁷⁷⁴ Katalinic, M., Z. Physik, 77, 257-270 (1932).

THE Konen, II., Ann. d. Physik (4), 9, 742-780 (1962).

THE Carter, H. D., and Campbell, A. N., Trans. Faraday Soc., 28, 479-496, 634-644 (1932) \rightarrow Trans. Electroch. Soc., 63, 419-423 (1933).

⁷⁷⁷ Shipley, J. W., Trans. Am. Electrochem. Soc., 55, 105-116 (1929).

⁷⁷⁸ Smith, H., Phil. Mag. (6), 27, 801-823 (1914).

⁷⁷⁹ Toriyama, Y., and Shinohara, U., Nature, 132, 240 (1933).

microsecond, the corona was white, and the illustration they give indicates that the spectrum was continuous.

Spark.

The spectra of sparks in liquids and between electrodes of various kinds have been much studied. A report on the subject, including a bibliography of 23 entries, has been published by J. A. Anderson. From such sparks, bubbles of gas are projected with considerable velocity, and in amounts greater than can be accounted for by electrolysis. For the water-spectrum of such sparks, see Section 46.

Y. Toriyama and U. Shinohara ⁷⁸¹ have found that there is no direct relation between the conductivity of a liquid dielectric and its break-down voltage: the break-down is an electronic phenomenon, the conduction, an ionic one. Using needle points separated by the distance d, and "impulse voltage chopped at the tail of the impulse wave," they found the following values for the break-down difference of potential (V) for water of conductivity 1.43×10^{-4} (ohm·cm)⁻¹:

These values of V were read from their graph.

53. MAGNETIC SUSCEPTIBILITY OF WATER

The permeability (μ) , the (volume) susceptibility (κ) , the specific susceptibility (χ) , and the density (ρ) of the medium are so related to the magnetic induction (B) and the resultant intensity (H) of the magnetic field that $B = \mu H = (1 + 4\pi\kappa)H$ and $\chi = \kappa/\rho$. The specific susceptibility (χ) is also called the coefficient of magnetization; if it is negative, the medium is said to be diamagnetic.

There is nothing to indicate that the value of χ for water depends upon the strength of the field. Fields of 1.2 to 40 kilogauss have been used.⁷⁸²

Detailed discussions of methods and of the several determinations of the susceptibility of water and its variation with the temperature have been published by P. Sève,⁷⁸³ A. Piccard,⁷⁸⁴ W. Johner,⁷⁸⁵ and recently, again by Sève.⁷⁸⁶

The most precise determinations yet made are those by A. Piccard and A. Devaud ⁷⁸⁷ giving $10^9\chi = -719.92 \pm 0.11$ cgsm $(10^9\kappa = -718.64)$ at 20 °C; and by H. Auer ⁷⁸⁸ giving $10^9\chi = -721.83 \pm 0.48$ cgsm at 20 °C.

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780 Anderson, J. A., Int. Crit. Tables, 5, 433 (1929).
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⁷⁸¹ Toriyama, Y., and Shinohara, U., Phys. Rev. (2), 51, 680 (L) (1937).

⁷⁸³ Hayes, H. C., *Phys. Rev.* (2), 3, 295-305 (1914); Wills, A. P., *Idem*, 20, 188-189 (1905); de Haas, W. J., and Drapier, P., *Ann. d. Physik* (4), 42, 673-684 (1913).

⁷⁸³ Sève. P., Ann. de chim. et phys. (8), 27, 189-244, 425-493 (1912) → Jour. de Phys. (5), 3, 8-29 (1913).

⁷⁸⁴ Piccard, A., Arch. sci. phys. ct nat. (4), 35, 209-231, 340-359, 458-482 (1913).

⁷⁸⁵ Johner, W., Helv. Phys. Acta, 4, 238-280 (1931) = Diss., Bern, 1930.

⁷⁸⁰ Sève, P., Congrès Internat. d'Elect., Sect. 2, Report 11, Paris, 1932.

⁷⁸⁷ Piccard, A., and Devaud, A., Arch. sci. phys. et nat. (5), 2, $455-485 \rightarrow 419$ (1920).

⁷⁸⁸ Auer, H., Ann. d. Physik (5), 18, 593-612 (1933).

The great difference between these two values (over 3 times the sum of their estimated uncertainties) is yet to be explained. The value accepted by the experts for the *International Critical Tables* is -720, essentially that obtained by Piccard and Devaud.

The surprising results reported by A. P. Wills and G. F. Boeker, 789 indicating a marked variability and a kind of hysteresis in the susceptibility of water, have not been confirmable, 788, 790 and seem to have been due to leaks in the apparatus.⁷⁰¹ But even after these have been eliminated the trend of the slope of the (χ, t) curve changes abruptly near 35 °C and near 55° C.791, 792

Cabrera and Fahlenbrach 790 found that the value of x for water from freshly melted ice was the same as that for water that had not recently

Table 185.—Specific Susceptibility of Water at 20 °C

It is believed that the first and the third entry represent the same observations. Itnit of wand um 1 ocem

	Unit of X and I	e=1 cgsm	
Year	Ref.ª	$-10^{9}\chi$	- 100 K
1912	Weiss and Piccard	719.3	720.6
1912	Sève	719.1 ^b	720.4
1913	Piccard	719.3	720.6
1913	de Haas and Drapier	721	722
1914	Ishiwara	720	721
1920	Piccard and Devaud	719.92 ± 0.11	721.19
1929	Int. Crit. Tables	720	721
1933	Auer	721.83 ± 0.48	723.11

a References:

Auer, H., Ann. d. Physik (5), 18, 593-612 (1933). de Haas, W. J., and Drapier, P., Ann. d. Physik (4), 42, 673-684 (1913).

Int. Crit. Tables. K. Honda, T. Ishiwara, T. Soné, and M. Yamada, Int. Crit. Tables. 6, 354, 356 (1929). Based on work of W. J. de Haas and P. Drapier, T. Ishiwara, A. Piccard and A. Devaud. 787, 9-160 P. Sève. 788, vol. 79 P. Weiss and A. Piccard, and A. P. Wills. Phys. Rev., 20, 188-189 (1905); references to the first two papers and to the next to the last are given elsewhere in this list.

Ishiwara, T., Sci. Rep. Tôhoku Imp. Univ. Sendai (1), 3, 303-319 (1914). Piccard, A., Arch. sci. phys. et nat. (4), 35, 209-231, 340-359, 458-482 (1913). Piccard, A., and Devaud, A., Idem (5), 2, 455-485 + 410 (1920).

Sève, P., Ann. de chim. et phys. (8), 27, 189-244, 425-493 (1912).

Weiss, P., and Piccard, A., Compt. rend., 155, 1234-1237 (1912).

This is the mean of the values for the two methods, as corrected by Piccard.

^b This is the mean of the values for the two methods, as corrected by Piccard loc. cit., and reduced to 20 °C by means of the coefficient 0.00012.

been frozen, the temperature being the same in both cases. But F. W. Gray and J. F. Cruikshank, 793 using a method by which the value of χ could be observed continuously, have reported that the numerical value of x for water from freshly melted ice increased for a time, reaching a maximum about 20 minutes after the melting, and then falling abruptly to a constant value. They seek to explain this variation in terms of the molecular structure proposed by Bernal and Fowler (p. 174).

⁷⁸⁰ Wills, A. P., and Boeker, G. F., Phys Rev (2), 42, 687-696 (1932).

⁷⁰⁰ Cabrera, B., and Fahlenbrach, H., Z. Physik, 82, 759-764 (1933).

⁷⁰¹ Wills, A. P., and Boeker, G. F., Phys. Rev. (2), 46, 907-909 (1934).

⁷⁹² Seely, S., Phys. Rev. (2), 52, 662 (L) (1937).

⁷⁹² Gray, F. W., and Cruikshank, . F., Nature, 135, 268-269 (L) (1935).

O. Specchia ⁷⁹⁴ has suggested that an interference method might be used to advantage for measuring the change in level upon which rest many of the determinations of χ .

The effect of variations in temperature upon the value of χ for water is so slight that it is difficult to measure. In the earlier measurements it was not only masked, but actually reversed, by errors and parasitic effects. In 1932 it seemed that $(\chi_t - \chi_{20})/\chi_{20} = \alpha(t-20)$ with $10^4\alpha = +1.31$, but later work indicates that the relation is not linear (see Table 186).

Explanations of the variation of χ with the temperature, generally in terms of changes in the polymerization, have been proposed by A. Piccard, ⁷⁸⁴, ⁷⁹⁵ W. Johner, ⁷⁸⁵ R. N. Mathur, ⁷⁹⁶ M. A. Azim, S. S. Bhatnagar, and R. N. Mathur, ⁷⁹⁷ B. Cabrera and H. Fahlenbrach, ⁷⁹⁸ G. Tammann, ⁷⁹⁹ and K. Honda and Y. Shimiza ⁸⁰⁰; and Mathur ⁷⁹⁶ has remarked that any progressive change in the uniformity with which the molecules are aligned by the field should result in a corresponding change in χ . From the observa-

Table 186.—Variation of the Specific Susceptibility of Water with the Temperature

From the data available in 1932 it appeared that the specific susceptibility of water varied linearly with the temperature, the most probable value of $\alpha \equiv (1/\chi_{20}) \cdot (d\chi/dt)$ being 0.000131 per 1 °C. But the observations by Piccard (P 1913)^a and by Mathur (Mr 1931)^a did not accord with that conclusion, and neither do the more recent observations by Auer (A 1933)^a, Cabrera and Fahlenbrach (CF 1934)^a, Wills and Boeker (WB 1934)^a, and Seely (See 1937)^a. The values tabulated by O. Specchia and G. Dascola ⁸⁰³ for ordinary water seem to be seriously in error.

The formula published by Cabrera and Duperier (CD 1924, 1925)^a and quoted in some compilations is now admitted to be wrong, actually defining a variation in the wrong direction; that given by Wills and Boeker (WB 1934)^a for the range 20 to 66 °C is $\chi/\chi_{20} = 1 - 1.3(t - 20)/10^4 - 0.7(t - 20)^2/10^6$, and defines the values given under WB in the second section of this table.

In the first section of the table are given the several values of the linear coefficient (α) that have been proposed and used. In the second are the several sets of values that have been obtained for $-\chi_{20}$ and for $-(\chi-\chi_{20})$. From them the corresponding values of χ for each of the tabulated values of t may be obtained; e.g., the observations of P give the values $-10^9(\chi-\chi_{20})=-0.9$ at 10° C and $-10^9\chi_{20}=719.3$, whence $-10^9\chi=719.3-0.9=718.4$ cgsm at 10° C.

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    Specchia, O., Atti. Accad. Naz d. Lincci (6), 7, 574-576 (1928).
    Piccard, A., Compt. rend., 155, 1497-1499 (1912).
    Mathur, R. N., Indian J. Phys., 6, 207-224 (1931).
    Azim, M. A., Bhatnagar, S. S. and Mathur, R. N., Phil. Mag. (7), 16, 580-593 (1933).
    Cabrera, B., and Fahlenbrach, H., Compt. rend., 197, 379-381 (1933).
    Tammann, G., Z. Physik, 91, 410-412 (1934).
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⁸⁰⁰ Honda, K., and Shimiza, Y., Sci. Rep. Tôhoku Imp. Univ. (Scudai), 25, 939-945 (1937).

Table 186—(Continued)

Unit of $\chi = 1$ egsm; of $\alpha = 1$ per 1 °C. Temp. = t °C

Linear coefficients. Not to be preferred.

Ref.a	\mathbf{WP}	Me	J, CJP	Sè	ABM	CF
Year	1912	1916	1930	1932	1933	1933
10 ⁴ α	1.2	1.00	1.31	1.31	1.39	1.15
ŧ	0 to 80	0 to 130	0 to 100		20 to 70	0 to 100

Various sets of values of χ_{20} and $\chi - \chi_{20}$. The Sè set has been computed by means of the linear coefficient $10^4\alpha = 1.31$ that was accepted in 1932; the others, except ABM, are experimental values, and do not vary linearly with the temperature; $-10^9 \chi = -10^9 \chi_{20} - 10^9 (\chi - \chi_{20})$.

$ \begin{array}{c} \operatorname{Ref} \stackrel{a}{\longrightarrow} \\ \operatorname{Year} \rightarrow \\ -10^{9} \chi_{20} \\ t \end{array} $	Sè 1932 720 ^b	P 1913 719.3	Me 1916 720b	Mr 1931 720b	ABM 1933 720 ^b 10 ⁹ (χ	A 1933 721.83	WB 1934 720 ^b	CF 1934 720 ^b	See 1937 720 ^b
0	-1.9	-2.0	-2.0		'A		-2.1	-1.66	-2.1
0 1 5 10	-1.8					-2.87			
5	-1.4	-1.5	0.0			- 2.01	-1.4	0.02	4.4
	-0.9	-0.9	-0.0	0.5		-1.16	-1.0	-0.83	-1.1
15	-0.5	-0.5		-0.5		-0.52	-0.5		_
20	0	0	0	0	0	0	0	0	0
25	+0.5	+0.3		+1.8		+0.41	+0.5	. 0.02	
30 35	0.9	0.8	+1.5	2 2	1	0.75 1.03	0.9	+0.83	+1.0
40	1.4 1.9	1.0 1.4	2.0	3.3	2	1.03	1.3 1.7	1.66	1.7
			2.0		4			1.00	1.7
45	2.4	1.7	2.4	5.1	2	1.54	2.0	0.40	2.2
50 55	2.8 3.3	$\frac{1.9}{2.2}$	3.1	7.2	3	1.78	$\substack{2.4\\2.7}$	2.48	2.2
60	3.8	$\frac{2.2}{2.4}$	3.1	1.2	4	2.23	2.9	3.31	3.0
65	4.2	2.6	3.1	8.4	-	2.20	3.2	0.01	0.0
70	4.7	2.7	2.0	0.4	5	2.71	3.4	4.14	4.4
75 75	5.2	2.8	3.9	9.7	3	2.71	0.4	4.14	5.2
80	5.6	2.9	4.4	9.1	6			4.97	5.2
85	6.1	3.0	7.7		U			1.7.	
90	6.6	3.0	5.7					5.80	
95	7.1	3.1	• • • •						
100	7.5	3.2	6.2						
110	8.5	0.2	7.2					7.06	
120	9.4		7.5					7.06	
130	10.4		7.7					7.06	

a References:

CD

Auer, H. 788

Azim, M. A., Bhatnagar, S. S., and Mathur, R. N., Phil. Mag. (7), 16, 580-593 (1933).

Cabrera, B., and Duperier, A. Ann. Soc. Esp. Fis. Quim., 22, 160-167 (1924); Jour. de Phys. (6), 6, 121-138 (1925).

Cabrera, B., and Fahlenbrach, H., Z. Physik, 82, 759-764 (1933); Ann. Soc. Esp. Fis. Quim., 31, 401-411 (1933); Z. Physik, 89, 166-178 (1934). See also, Ann. Soc. Esp. Fis. Quim., 32, 525-537. 538-542 (1934).

Cabrera, B., Johner, W., and Piccard, A., Compt. rend., 191, 589-591 (1930). Johner, W. 785

Marke, A. W., Fortsch. Physik, 73-, 74 (1918) — Overs. K. Danske Vid. Sclsk. Forh. (Kopenhagen), 5 and 6, 395-413 (1916).

Mathur, R. N. 786

Seely, P. 786

Seely, S., Phys. Rev. (2), 52, 662 (L) (1937). CF

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Seely, S., Phys. Rev. (2), 52, 662 (L) (1937). Wills, A. P., and Boeker, G. F.⁷⁰² Weiss, P., and Piccard, A., Compt. rend., 155, 1234-1237 (1912).

Assumed by the compiler in deriving the values of $(\chi - \chi_{20})$.

tions of Cabrera and Fahlenbrach and Rao's estimate of the relative abundance of the several polymers,⁸⁰¹ L. Sibaiya ⁸⁰² has computed the value of χ for each of the polymers assumed by Rao, finding: $10^9\chi_{20} = -775.5$ for (H_2O) , -722.2 for $(H_2O)_2$, and -701.3 for $(H_2O)_3$.

54. VERDET CONSTANT OF WATER

When plane-polarized light passes a distance l through a substance in a uniform magnetic field of strength II, the angle between the direction of H and the direction of advance of the light being θ , then the plane of polarization is rotated through an angle α , in the direction in which a right-handed screw lying along l must be turned in order that its advance shall be in the same direction as that of the component along l of the light, such that $\alpha = VlH\cos\theta$, V being a factor determined by the substance, its temperature, and the wave-length of the light. If V is positive and θ is zero, the rotation is in the direction of the amperian currents that are equivalent to the field H.

This phenomenon is often called the Faraday effect, and the factor V is commonly called the Verdet constant.

The effect appears very quickly after the field is applied. The exact amount of lag, if any, is not known. Not only does the early conclusion of J. W. Beams and F. Allison 804 that the lag for water exceeds that for CS₂ by 1.1 mµsec seem to be incorrect as to the numerical value (cf. F. Allison 805), but doubt has even been cast upon their interpretation of their observations (see, e.g., J. W. Beams and E. O. Lawrence, 806 E. Gaviola, 807 F. G. Slack, R. L. Reeves, and J. A. Peoples, Jr. 808). There is at present no generally accepted experimental evidence of any lag at all. Similar remarks apply to Allison's early conclusions that the difference between the lags for water and for CS₂ vanishes when the liquids are exposed to x-rays, and that such exposure increases slightly the value of the Verdet constant. 808a

Effect of Temperature.

For the D-lines ($\lambda=0.5893~\mu$) and within the range 4 °C to 97.7 °C, $10^4V_t=131.1-0.00400t-0.000400t^2$ minute-of-arc per cm·gauss.⁸⁰⁹ This formula is equivalent to that given by Λ . Cotton and R. Lucas,⁸¹⁰ and leads to the following values:

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802 Sıbaiya, L., Current Sci., 3, 421-422 (1935).
803 Speechia, O., and Dascola, G., Nuovo Cim. (N. S.), 12, 606-609 (1935).
804 Beams, J. W., and Allison, F., Phys. Rev. (2), 29, 161-164 (1927).
805 Allison, F., Idem, 30, 66-70 (1927).
806 Beams, J. W., and Lawrence, E. O., J. Franklin Inst., 206, 169-179 (1928).
807 Gaviola, E., Phys. Rev. (2), 33, 1023-1034 (1929).
808 Slack, F. G., Reeves, R. L., and Peoples, J. A., Jr., Phys. Rev. (2), 46, 724-727 (1934).
808 Allison, F., Phys. Rev. (2), 31, 158-159 (A) (1928).
808 Rodger, J. W., and Watson, W., Phil. Trans. (A), 186, 621-655 (1895), → Z. physik. Chem.,
19, 322-363 (1896).
810 Cotton, A., and Lucas, R., Int. Crit. Tables, 6, 425 (1929).
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$$t$$
 0 20 40 60 80 100 °C 10 4V_D 131.1 130.8 $_8$ 130.3 $_0$ 129.4 $_2$ 128.2 $_2$ 126.7 $_0$

For water, the ratio of V_D to the density is essentially constant if t < 20 °C, but above 20 °C the ratio increases almost linearly with t, about $1.4' \times 10^{-6} \text{cm}^{-1} \text{ gauss}^{-1} \text{ cm}^3/\text{g} \cdot \text{deg.}^{809}$.

It is generally assumed, and the data available in 1932 indicated (e.g., F. Schwers ⁸¹¹), that V_{λ}/V_{D} is essentially independent of the common temperature, at least over a moderate range including 20 °C.

But the recent observations by F. G. Slack, R. L. Reeves, and J. A. Peoples, Jr. ⁸¹² suggest that V_{λ}/V_{D} may vary slowly with the temperature; they report as follows, V_{546} being the value of V for $\lambda = 0.546 \,\mu$.

Their value of V_D for t = 20 °C being 0.77 per cent greater than the one generally accepted (see Table 187).

Dispersion of the Verdet Constant.

As just stated, $(V_{\lambda}/V_D)_t$ is essentially independent of t if t is not far from 20 °C, but the several sets of measurements of V for a fixed temperature and various wave-lengths exhibit annoying discrepancies, and a direct graphical comparison of them is not satisfactorily accurate. Furthermore, it is quite laborious to determine the deviation of each value from that demanded either by the equation (1) proposed by P. Joubin 813

$$V\lambda = 0.002788_5(n - 77.65\lambda dn/d\lambda) \tag{1}$$

or by that (2) given by S. S. Richardson 814 *

$$n\lambda^2 V = 0.003265_5 \left[\left(\frac{\lambda^2}{\lambda^2 - 0.01891} \right)^2 + 0.7381_0 \right]$$
 (2)

or by that (3) used by U. Meyer 815

* In International Critical Tables, 6, 425, where the expression in parentheses is written in the form $\left(\frac{\lambda^2}{\lambda^2-\lambda_1^2}\right)^2$, λ_1 is incorrectly given as 1260.4A. It should be 1375.1A, as Richardson gives $\lambda_1^2=0.01891~\mu^2$

Exactly similar formulas, but with different constants, have been proposed by G. Bruhat and A. Guinier, 816 $n\lambda^2V=0.0039470$ [0.43090 + $\lambda^4/(\lambda^2-0.01680)^2$], and by I. T. Pierce and R. W. Roberts, 617 $n\lambda^2V=0.0038723$ [0.45658 + $\lambda^4/(\lambda^2-0.01719)^2$], as representative of their own sets of observations. Each thinks that the formula given defines V to within 1 in 1000; the first, for the visible and ultraviolet spectrum; the second for the infrared.

- 811 Schwers, F., Bull. Acad. Roy. Bela., 1912, 719-752 (1912).
- 812 Slack, F. G., Reeves, R. L., and Peoples, J. A., Jr., Phys. Rev. (2), 46, 724-727 (1934).
- 813 Joubin, P., Ann. de chim. et phys. (6), 16, 78-144 (1889).
- 814 Richardson, S. S., Phil. Mag. (6), 31, 232-256 (1916).
- 815 Meyer, U., Ann. d. Physik (4), 30, 607-630 (1909).
- 816 Bruhat, G., and Guinier, A., Jour. de Phys. (7), 4, 691-714 (1933).
- 817 Pierce, I. T., and Roberts, R. W., Phil. Mag. (7), 21, 164-176 (1936).

Table 187.—The Verdet Constant of Water

(See Figs. 7 and 8)

For the D-lines ($\lambda = 0.5893 \,\mu$) the generally accepted value of V for water at 20 °C is $V_{20} = 0'.01309$ per cm-gauss = 3.808 microradians/cm-gauss, which essentially agrees with that (0.01308) given by A. Cotton and R. Lucas. 818

 V_n is the norm computed by means of the arbitrary formula (4,5) constructed so as to represent closely the observations of Mi and of I (see text), the required values of Δ being those here given. Except the Ri values, computed by means of the equation as given by Richardson, each set of values has been so adjusted by a flat percentile correction, as to give V = 0'.01309 for the D-lines.

Examples: For 20 °C and $\lambda = 0.2428 \,\mu$, Ri's equation gives $10^4 V = 1159.8 + 5.6 = 1165.4$; for 20 °C and $\lambda = 0.2482 \,\mu$, the value found by Mi was 1084, by Ro 1086, and by BG 1079, none of these giving the fifth digit.

Unit of $\lambda = 1$ $\mu = 10^{-4}$ cm; of V and $\Delta = 1'$ per cm-gauss. Temp. = t °C

I. Best values for the D-lines ($\lambda = 0.5893$).

t	V_t	10^4V_{20}	References ^a
20	0.01309	130.9	Rodger and Watson (1895)
18	0.01309	130.9	Agerer (1905)
20	0.01309	130.9	Richardson (1916)
17	0.01306	130.7	Stephens and Evans (1927)

II. The better values throughout the spectrum. Temp. = $20 \,^{\circ}$ C.

λ	Referencesa 1042	→ 10V _n	Mi	Ri^b	Ro	$V - V_n$	L	v.S
0.2428	+22.8	1159.8		+ 5.6	10-(, - , n,		
82	15.2	1079.6	+4	1 3.0	+6	- 1		
96	13.6	1060.2	, -		, 0	-	- 12	
0.2536	10.0	1008.5	0					
37	9.8	1007.2			+10			
40	9.6	1003.8				+1		
76	7.2	961.9	-4					
0.2652	3.1	881.3				+1.8		
55	2.9	878.3	+2		+12			
0.2700	+0.8	835.8	0					
50	-1.1	792.4					- 12	
53	-1.1	790.0	0					
0.2804	-2.5	749.5	0			+3.4		
05	-2.6	748.7			+7			
94	-4.0	686.7	0					
0.2925	-4.3	667.0	- 1					
68	-4.6	641.3	0					
0.3023	-4.7	610.9	0	. 1.6				
34	-4.8	605.0		+1.6				
0.3100	-4.8	571.7					- 10	
26	-4.8	559.7	-1					
30	-4.8 -4.8	558.0			. 0	+2.9		
31 32	-4.8 -4.8	557.5	- 1		+8			
		557.1	1					
0.3303	-4.9 -4.9	486.1		+1.2	. =	. 0 7		
41	-4.9	472.4 472.0	0		+5	+2.7	,	
42	-4.9	4/4.0	U					

⁸¹⁸ Cotton, A., and Lucas, R., Int. Crit. Tables, 6, 425 (1929).

Table 187—(Continued)

	References"	1047	Mi	Rib	Ro	BG	L	v.S
λ 0.3580 0.3609	10 ⁴ Δ - 4.4 - 4.4	10 ¹ V _n 398.5 390.7			104($V - \overline{V}_n$) ——	-5	-5.1
11 12	-4.4 -4.4	390.1 389.9		+1.2			- 3	
52	-4.4 -4.2 -4.2	379.8	-1	+1.3				
55 63	-4.2	379.1 377.0	-1			+2.4		
65 0.3729	$-4.1 \\ -3.9$	376.6 361.6			+5			-4.0
0.3886 0.3907	$-3.4 \\ -3.4$	328.0 323.8	- 2				-3	
62 0.4046	-3.2 -3.0	313.5 297.2		+1.0			-2	
47 54	-3.0 -3.0	298.4 297.3	0		+3	+1.6	_	
78 0.4199	- 2.9 - 2.6	293.2 274.3	- 1				-4	
0.4307 08	-2.3 -2.3	258.9 258.8					-4	-2.2
40 41	-2.3 -2.2 -2.2	254.5 254.4		$^{+0.7}_{+0.7}$			-4	
58	-2.2	252.1	6	70.7	0	+1.3		
0.4400	-2.1 -2.0	252.1 246.9	$-\frac{1}{0}$					
05 0.4505	-2.0 -1.8	246.3 234.1	+0.8				-3 -3	
29 0.4605	- 1.8 - 1.6	232.4 223.0	+0.5				-3	
78 0.4705	-1.4 -1.3	215.4 212.7	+0.6	+0.5				
0.4805 61	$-1.1 \\ -1.0$	203.2 198.2	+0.7					-3.2
0.4905 16	-0.9 -0.9	194.2 193.3	0.0		+0.1			
21 ° 58	-0.9 -0.8	192.8 189.8		+0.2			-4	
0.5005 0.5105	-0.8 -0.6	185.8 178.1	0.0 0.0					
0.5210	-0.5 -0.4	170.4 166.2	-0.1					-1.5
0.5310 0.5410	-0.4 -0.3	163.6 157.1	$^{+0.1}_{-0.2}$					0
61°,d 0,5515	-0.3	154.0 150.8	$+1 \\ +0.2$	+0.1	-0.6			
0.5615 0.5715	$ \begin{array}{r} -0.2 \\ -0.2 \\ -0.1 \end{array} $	145.0 139.7	$+0.1 \\ -0.3$		-	Ia -		-
80°	-0.1	136.4			-0.9		1	-
0.5815 0.5893^d	0	134.7 130.9	-0.3	0.1	0.0	0.0		0.0
0.5920 0.6000	0	129.6 125.9	+0.2	•			0	
20 0.6104	0 +0.1	125.1 121.5	+0.3		+0.3			
20 0.6220	0.1 0.1	120.8 116.7	$+0.1 \\ +0.4$					
0.6320	0.1	112.8	+0.5					

Table 187—(Continued)

					-			
λ	References ^a -	→ 1C4V'n	Mi	Rıb	Ro 104(V	BG	L	v.S
0.6420	0.2	109.2	+0.3					
0.6530	0.2	105.4	+0.4					
63	0.2	104.3						-0.6
0.6708	0.22	99.6			+0.2			•
0.7000	0.26	91.2				-0.2		
65	0.26	89.4		0.0				
0.8000	+0.3	69.0				-1.7	+1	
80	0.3	67.7		0.0				
71	0.4	58.0		0.0				
0.900	0.4	54.3				-3.4		
1.000	0.4	43.8				-3.0	0	
28	0.4	41.4		0.0				
1.100	0.4	33.6				-0.4		
1.200	0.3	30.2				-1.4		
50	0.3	27.8				3.4	+1	
56	0.3	27.6		0.0				
1.300	0.2	25.6		3.0		+0.7		
11000	0.2	20.0				1 0.7		

a References:

Agerer, F., Sitz. Akad. Wiss. Wicn (Abt. IIa), 114, 803-830 (1905).
Bruhat, G., Jour. de Phys. (7), 5, 152 (1934) & Bruhat. G., and Guinier, A., Idem, 4, 691-714 (1933) > Compt. rend., 197, 1028-1030 (1933).
Ingersoll, L. R., Phys. Rev., 23, 489-497 (1906).
Ingersoll, L. R., J. Opt. Soc. Amer., 6, 663-681 (1922).
Landau, St., Physik. Z., 9, 417-431 (1908).
Miescher, E., Helv. Phys. Acta, 3, 93-133 (1930); 4, 398-408 (1931).
Richardson, S. S. R. W., Phil. Mag. (7), 9, 361-390 (1930).
Rodger, J. W., and Watson, W. 1909.
Stephens, D. J., and Evans, E. J., Phil. Mag. (7), 3, 546-565 (1927).
von Schaik, W. C. L., Arch. Néerl. des Sci. Exact et Nat., 17, 372-390 (1882); 21,

Μi

Ro RW

These values are those defined by Richardson's equation; his observed values are as follows:

 $^{\circ}R.$ de Mallemann, P. Gabiano, and F. Suhner 818n have reported the following values, reduced from 11.5 $^{\circ}C$ to 20 $^{\circ}C$ by the compiler:

$$\lambda$$
 0.436 0.492 0.546 0.578 μ 104V 254.1 194.8 154.2 136.4

 4 F. G. Slack, R. L. Reeves, and J. A. Peoples, Jr. 812 have reported the following high values for 20 °C: $10^{4}V=156$ for $\lambda=0.5461,$ and 131.9 for $\lambda=0.5893~\mu$.

$$nV = 0.005606\lambda^2/(\lambda^2 - 0.013253)^2 \tag{3}$$

 $(n = index of refraction; \lambda = wave-length, unit = 1 \mu$; the first constant in each equation has been so chosen as to make $V_D = 0'.01309$ per cm-gauss.)

Consequently, a simpler expression (4) has been adopted as a norm with which to compare both the observations and the proposed equations

$$V_n - \Delta = \frac{4.2347}{1000\lambda^2} \left(\frac{\lambda^2}{\lambda^2 - 0.012097} \right)^2 \tag{4}$$

or

818a de Mallemann, R., Gabiano, P., and Suhner, F., Compt. rend., 202, 837-838 (1936)—Gabiano, P., Jour. de Phys. (7), 7, 84S (1936).

$$(V_n - \Delta)^{-0.5} = 15.367\lambda - \frac{0.18589}{\lambda}$$
 (5)

The values of Δ (see Table 187), varying slowly and continuously with λ , have been so chosen as to make V_n approximately represent the observations of Miescher (Mi of Table 187) and of Ingersoll (I and I' of Table 187). From the value of $(V_n - \Delta)^{-0.5}$ that of $V_n - \Delta$ can be directly obtained by the use of Barlow's Tables of Squares, etc., either by entering

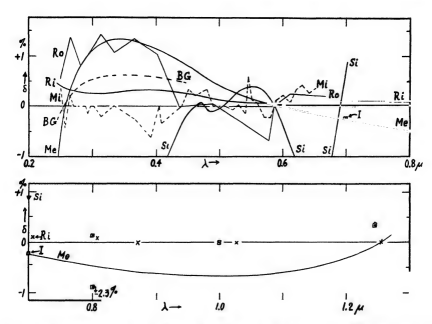


FIGURE 7. Deviations of the Observed and Computed Values of the Verdet Constant for Water from those Defined by Formula (4).

 $\delta=(V-V_n)/V_n$, where V_n is the value defined by the arbitrarily chosen norm, formula (4); $\lambda=$ wave-length. Unit of $\delta=0.01$; of $\lambda=1~\mu=0.0001$ cm.

Each set of data, except Richardson's curve (Ri), has been multiplied by such a constant as to make $\delta=0$ for the D-lines ($\lambda=0.5893~\mu$), but in that region the Me-curve has accidentally been drawn a little too high, not enough to be of real significance. In the upper section of the figure ($\lambda<0.8~\mu$), the Ri and Me curves represent, respectively, the formulas given by Ri and by Me; the values given by Si are said to have been read by him from a smoothing curve, and define the curve here given; all the BG-values lie within 0.1 per cent of the curve so marked. The successive individual determinations by Ro, also those by Mi, are connected by straight lines, and those by I are indicated by squares. In the lower section, the Me-curve is continued, the computed Ri-values are indicated by crosses, the I-values by squares, and the Si-value by an inverted triangle. One of the I-values near $\lambda=0.8$ belongs far below the boundary of the figure, at -2.3 per cent, as indicated.

References:

as indicated.

- Bruhat, G., Jour. de Phys. (7), 5, 152 (1934) ← Bruhat and Guiner, Idem, 4, 691-BG 714 (1933).
- Ingersoll, L. R., Phys. Rev., 23, 489-497 (1906); J. Opt. Soc. Amer., 6, 663-681 Ι
- (1922).
 Meyer, U., Ann. d. Physik (4), 30, 607-630 (1909).
 Miescher, E., Helv. Phys. Acta, 3, 93-133 (1930); 4, 398-408 (1931).
 Richardson, S. S., Phil. Mag. (6), 31, 232-256 (1916).
 Roberts, R. W., Idem (7), 9, 361-390 (1930).
 Siertsema, L. H., Arch. Néerl. des Sci. (2), 6, 825-833 (1901). Me
- Mi Ri
- Ro

the column of \sqrt{n} and taking the corresponding number in the column of 1/n, or by entering the column of 1/n and taking the corresponding number in the column of n^2 .

The various deviations from V_n are shown in Figs. 7 and 8, and those of the better series of observations are given in Table 187 in such a way that the individual observations in those series may be recovered if desired. It will be noticed from the graphs that Joubin's equation, though commonly included in compilations of data, is entirely unsatisfactory; whether the same is true of Meyer's is not clear, as that does approximately represent Roberts' observations.

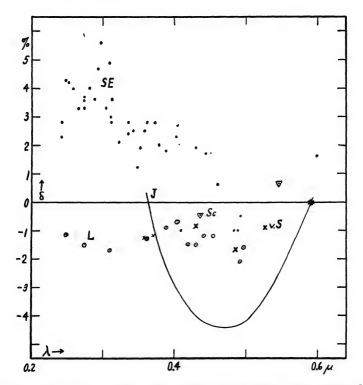


FIGURE 8. Deviations of Other Observed and Computed Values of the Verdet Constant of Water from those Defined by Formula (4).

 $\delta = (V - V_n)/V_n$, where V_n is the value defined by the arbitrarily chosen norm, formula (4); $\lambda =$ wave-length. Unit of $\delta = 0.01$; of $\lambda = 1$ $\mu = 0.0001$ cm.

This differs from Fig. 7 both in scale and in data. The computed J-curve is obviously unsatisfactory. D-lines). Its constant was so chosen as to make the curve pass through the star ($\delta = 0$ for the

References:

Joubin, P., Ann. chim. phys. (6), 16, 78-144 (1889). (circles) St. Landau, Physik. Z., 9, 417-431 (1908). (inverted triangles) Schwers, F., Bull. Acad. Roy. de Belg., 1912, 719-752 (1912). (dots) Stephens, D. J., and Evans, E. J., Phil. Mag. (7), 3, 546-565 (1927). (crosses) van Schaik, W. C. L., Arch. Néerl. des Sci., 17, 372-390 (1882); 21, 406-431 (1887).

55. Magnetic Birefringence of Water

In a magnetic field, water is negatively birefringent, its index of refraction (n_p) for light in which the electric vector is parallel to the field is less than that (n_t) for light in which that vector is transverse to the field. If λ = wave-length of the light in a vacuum, and H is the strength of the magnetic field, then the coefficient (C_m) of magnetic birefringence (sometimes called the Cotton-Mouton constant) is defined by the equation $C_m = (n_p - n_t)/\lambda H^2$.

For water, C_m is very small, of the order of one-thousandth of the value for nitrobenzene. M. A. Haque ⁸¹⁹ found $10^{14}C_m = -0.3_0$ cm⁻¹ gauss⁻², and S. W. Chinchalkar⁸²⁰ and H. A. Boorse ⁸²¹ each found -0.3_7 . An earlier, less accurate, measurement by M. Ramanadham ⁸²² gave -1.1, nearly 3 times the values found by the others. A. Cotton and T. Belling ⁸²³ have reported -0.14, about $\frac{1}{3}$ of the value found by Haque.

IIC. ICE

56. Foreword

Of the several treatises dealing with ice, the one that seems to be by far the most comprehensive that has come to the compiler's attention is that by A. B. Dobrowlski: "Historja Naturalna Lodu" ("The Natural History of Ice"). The manuscript was completed in 1916, but the volume was not published until 1923. It is printed in Polish (a language not read by the compiler) and contains 940 pages, including a French translation of the introduction and table of contents, and an author index of over 1000 names. Its purpose and scope are thus defined: "L'Histoire Naturelle de la Glace est un essai de synthèse des recherches faites, dans la nature et dans le laboratoire, sur la glace de tout aspect et de toute origine. C'est une sorte d'index de tous les problèmes relatifs à ce corps si important, et si peu connu encore, avec un exposé de l'histoire de chacun de ces problèmes, des résultats acquis, des questions litigieuses et des lacunes."

57. Types of Ice

For crystallographic forms of ice, see Section 59; for x-ray studies of ice, see Sections 60 and 74.

Besides ordinary ice, more particularly designated as ice-I, and vitreous ice obtainable at low temperatures, six other distinct varieties of ice, each having a definite region of stability, are known ¹; two forms of ice-I have been reported ²; and ice that is denser than water, though formed at a

⁸¹⁹ Haque, M. A., Compt. rend., 190, 789-790 (1930).

⁸²⁰ Chinchalkar, S. W., Indian J. Phys., 6, 165-179 (1931).

⁸²¹ Boorse, H. A., Phys. Rev. (2), 46, 187-195 (1934).

⁸²² Ramanadham, M., Indian J. Phys., 4, 15-38 (1929).

Cotton, A., and Belling, T., Compt. rend., 198, 1889-1893 (1934).
 Bridgman, P. W., Proc. Amer. Acad. Arts Sci., 47, 439-558 (1912); J. Chem'l Phys., 3, 597-605 (1935); 5, 964-966 (1937).

² Seljakov, N. J., Compt. rend. Acad. Sci. URSS, 10, 293-294 (1936); 14, 181-186 (1937).

pressure of less than 1 atm, has been reported twice, but in neither case could its formation be repeated. Shaw states that it is denser than water; and Cox says it "sank slowly to the bottom and remained there with, perhaps, one third of an inch of clear water above it." (A few rather casual observations by the compiler have suggested that these conclusions may rest upon an illusion. The capillary pull of the gas-liquid surface between the walls of the bulb and the ice will depress the ice, and if the surface of the ice is concave, the concavity may become filled with water, producing a striking impression that the ice is fully submerged. The requirements being rather exacting, their accidental fulfillment will not be frequent.) Evidence for the existence of several unstable types of ice was presented by G. Tammann, but not generally accepted; and nothing has been heard of those unstable types for many years.4 In particular, his claims for the one he called ice-IV led to a discussion with Bridgman,4 and resulted in Bridgman's leaving that designation open when he published his 1912 paper; but now he has adopted it as the designation of an ice recently discovered.⁵ Bridgman's ice-IV should not be confused with Tammann's supposed ice of the same designation.

Ignoring for the present the vitreous and unstable types, and the dense ice reported from Canada, no type except the familiar variety, specifically denoted as ice-I, can exist under pressures much less than 2000 atm unless the temperature is very low, but at low temperatures the transformation of both ice-II and ice-III to ice-I proceeds so slowly, even at atmospheric pressure, that Tammann ⁶ has succeeded in removing them from the pressure chamber and examining them. He found ice-III to be a colorless, pellucid aggregate of coarse crystallites, which slowly swelled and broke up into a coarse white powder. In contact with warm objects it acquired a porcelain-like appearance.⁷

It has been said that vitreous ice is formed when small drops of water are quickly chilled to a low temperature, say to $-12\,^{\circ}\text{C}$ or lower.⁸ Beilby stated that this "ice" is perfectly transparent, shows under a microscope no evidence of crystalline structure, but at once crystallizes throughout when mechanically strained by a light pressure with a polished steel burnisher. Hawkes ⁸ regarded it as merely supercooled water, but that conflicts with the observations of H. C. Sorby, ⁹ L. Dufour, ¹⁰ and others, who have observed that water remains fluid even at temperatures ranging from $-12\,^{\circ}\text{C}$ to $-20\,^{\circ}\text{C}$. Sorby is especially definite in his statement of its fluidity. Using volumes of several cubic centimeters, the compiler has noticed that water at $-20\,^{\circ}\text{C}$ appears to the eye to be as fluid as it is at

⁸ Cox, J., Trans. Roy. Soc. Canada, Sect. III (2), 10, 3-4 (1904); Shaw, A. N., Idem (3), 18, 187-189 (1924).

⁴ See Tammann, G., Ann. d. Physik (4), 2, 1-31 (1900); Z. physik. Chem., 84, 257-292 (1913); 88, 57-62 (1914); Bridgman, P. W., Proc. Amer. Acad. Arts Sci., 47, 439 558 (1912); Z. physik. Chem., 86, 513-524 (1913); 89, 252-253 (1915).

⁸ Bridgman, P. W., J. Chem'l Phys., 3, 597-605 (1935).

⁶ Tammann, G., "The States of Aggregation," New York, 1925; Z. anorg. Chem., 63, 285-305 (1909). Cf. Bridgman, P. W., J. Franklin Inst., 177, 315-332 (1914).

room temperature. Although a vitreous solid may be an undercooled liquid of great viscosity, there seems to be no valid reason for assuming, as is frequently done, that every vitreous solid is of that nature; unless, of course, the term liquid is so defined as to cover everything that is neither gaseous nor crystalline. F. Simon¹¹ has presented reasons and experimental data in support of the idea that there are in reality two types of vitreous "solid." One is a true glass and the other is a supercooled liquid. the latter being in internal thermodynamic equilibrium, and the former not. Nevertheless, in view of the work now to be mentioned, it seems probable that Beilby's observations are in some way erroneous. The work should be repeated.

Using larger volumes of pure water, the late E. W. Washburn of the National Bureau of Standards failed to obtain vitreous ice even when the water was quickly chilled with liquid air, but when the viscosity was increased by the addition of a little sugar (about 1 per cent) vitreous ice was obtained¹². And E. F. Burton and W. F. Oliver ¹³ reported that when water-vapor condenses on copper at -110 °C, or lower, the resulting ice is vitreous, and as the temperature is then raised, say to -50 °C, the ice gradually crystallizes. This same phenomenon-vapor being condensed to an amorphous solid on a cold surface, and that solid becoming crystalline when the temperature is increased—had been previously reported by L. R. Ingersoll and S. S. DeVinney 14 for the case of nickel sputtered in hydrogen. The ice formed by condensation at -80 °C has the normal crystalline arrangement; as the temperature of formation is lowered, the crystal structure becomes less regular.

Of the known varieties of crystalline ice, all except the familiar one (ice-I) are denser than water under the same conditions of temperature and pressure. One consequence of this is that the pressure that is exerted by the freezing of water in a confined space can under no circumstance greatly exceed 2000 kg*/cm² (say 30,000 lb*/sq. in.), because the bulky ice-I cannot exist under such pressures. See phase diagram, Section 93, or H. T. Barnes.15

In what follows, we shall confine our attention, unless the contrary is clearly indicated, to the familiar variety of ice (ice-I), that which melts at 0 °C when under a pressure of 1 atm.

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7 Cf. Dewar, J., Chem. News, 91, 216-219 (1905).
<sup>8</sup> Beilby, G., "Aggregation and Flow of Solids," p. 195+, London, 1921; Hawkes, L., Nature, 123, 244 (1929).
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⁹ Sorby, H. C., Phil. Mag. (4), 18, 105-108 (1859). 10 Dufour, L., Ann. d. Physik (Pogg.), 114, 530-554 (1861).

¹¹ Simon, F., Z. anorg. allgem. Chem., 203, 219-227 (1932); Trans. Faraday Soc., 33, 65-73 (1937).

¹² Washburn, E. W., Oral communication, 1933. Work not published.

¹⁸ Burton, E. F., and Oliver, W. F., Proc. Roy. Soc. (London) (A), 153, 166-172 (1935); →Nature, 135, 505-506 (L) (1935).

¹⁴ Ingersoil, L. R., and DeVinney, S. S., Phys. Rev. (2), 26, 86-91 (1925).

¹⁵ Barnes, H. T., "Ice Engineering," p. 91, Montreal, Renouf Publishing Co., 1928.

58. APPEARANCE OF ICE-1

Ordinary ice, Ice-I, is crystalline, and has a conchoidal fracture and a vitreous luster. In large masses, it is vividly blue, owing to the scattering of light by its large molecules.^{15, pp. 8, 9, 18} It has long been held that water contains ice in solution, the amount increasing as the temperature decreases (see p. 164). This causes the color of water to change as the temperature falls. Barnes says: "It is a remarkable sight in winter to watch the varying shades of the river water as the temperature changes. Just at the freezing point the color changes rapidly and old river men can tell the approach of the ice forming period by the color." ^{15, p. 10}

59. Forms and Formation of Ice

All interested in this subject should read the very interesting and beautifully illustrated publications by W. A. Bentley ¹⁶ and by W. A. Bentley and W. J. Humphreys. ¹⁷ They are much broader than the titles indicate, most of the topics considered in this section being discussed.

The information contained in this section is almost entirely descriptive, and has been arranged under the following heads:

Crystallographic structure Production of homogeneous ice

Structure of ice Monocrystals

Internal melting Freezing of supercooled water

Flowers of ice Icicles
Formation of frazil, or needle, ice Hail

Formation of an ice-sheet Snow and frost

Growth and orientation of crystals Glaciers Recrystallization Sea-ice

Regelation Icebergs (see Glaciers, Sea-ice)

Purity

Crystallographic Structure.

(For ratio of axes and for fine-structure, as revealed by x-rays, see Section 60.)

X-ray examination indicates that ice-II is characterized by a side-centered orthorhombic cell containing 8 molecules, ¹⁸ and ice-III by a body-centered orthorhombic cell containing 16 molecules and having a:b:c = 1.73:1:1.22.¹⁹ Nothing is known of the crystallographic structure of any of the other ices except ice-I, to which we now turn.

The published data relative to the crystallographic structure of ice-I are confused by changing nomenclature, and are otherwise conflicting. The

¹⁶ Bentley, W. A., Monthly Weather Rev., 29, 212-214 (1901); 35, 348-352, 397-403, 439-444, 512-516, 584-585 (1907).

¹⁷ Bentley, W. A., and Humphreys, W. J., "Snow Crystals," New York, McGraw-Hill Book Co., 1931.

¹⁸ McFarlan, R. L., J. Chem'l Phys., 4, 60-64 (1936) → Phys. Rev. (2), 49, 199 (A) (1936).

¹⁹ McFarlan, R. L., Idem, 4, 253-259 (1936) → Idem, 49, 644 (A) (1936).

subject is further complicated by recent observations indicating that ice-I can exist in either of two forms: α -ice, which is hexagonal and appears when water near 0 °C freezes, and β -ice, which is rhombohedral and appears when water that is supercooled by at least a few degrees freezes.²⁰

From a consideration of all observations prior to 1906, P. H. Groth ²¹ thought it probable that ice-I belongs in the ditrigonal-pyramidal class of the trigonal system; A. E. H. Tutton ²² thought that more recent work, such as that of F. Rinne, ²³ shows that it belongs in the hexagonal bipyramidal class of the hexagonal system. The subject has been reviewed still more recently by A. B. Dobrowolski, ²⁴ who has concluded from his study of thousands of ice crystals that the symmetry of ice is that of the ditrigonal-pyramidal class of the trigonal system.

- W. Althorg and W. Troschin²⁵ have described some unusual forms of ice found in the ice cave near Kungur in the Ural mountains; and G. Tammann and K. L. Dreyer ²⁶ have given a popular account of the freezing of water and of some of the peculiarities of artificial ice.
- E. S. Dana ²⁷ states that ice crystallizes in the hexagonal system. This is accepted by H. T. Barnes, ²⁸ who states that the crystals are probably hemimorphic, that the crystal faces are rarely distinct, and that the crystals are hard to measure.
- L. J. Spencer ²⁹ assigned ice to the holosymmetric class of the rhombohedral division of the hexagonal system. This probably harmonizes the observations of the early observers, who variously reported that the crystals were hexagonal and rhombohedral; in some cases, the latter term seems to have been used as an equivalent of the former. That the primary form was the rhombohedron in the cases reported by H. Abich, ³⁰ by Sir David Brewster, ³¹ and by E. D. Clarke ³² seems beyond question; but the assertion of F. Leydolt ³³ that in his extended study of ice from various sources, he had found only rhombohedric forms, possibly means no more than that they belonged to the hexagonal system, without distinction between the rhombohedral and the hexagonal classes of that system. Likewise the many

²⁰ Seljakov, N. J., Compt. rend. Acad. Sci. URSS, 10, 293-294 (1936); 11, 227 (1936); 14, 181-186 (1937).

a Groth, P. H., "Chemische Krystallographie," Vol. 1, p. 66 (1906).

²² Tutton, A. E. H., "Crystallography and Practical Crystal Measurement," Vol. 1, p. 543 (1922).

²⁸ Rinne, F., Ber. Sächs. Ges. Wiss. (Math.-Phys.), 69, 57-62 (1917).

²⁴ Dobrowolski, A. B., Bull. Soc. Fr. Mineral., 56, 335-346 (1933).

²⁵ Altberg, W., and Troschin, W., Naturwissenschaften, 19, 162-164 (1931).

²⁶ Tammann, G., and Dreyer, K. L., Idem, 22, 613-614 (1934).

²⁷ Dana, E. S., "A Textbook of Mineralogy," 3rd ed., p. 411, revised by W. E. Ford, New York, John Wiley & Sons, 1922.

²⁸ Barnes, H. T., "Ice Formation," p. 74, New York, John Wiley & Sons, 1906; "Ice Engineering," p. 18,

^{**}Spencer, L. J., 'Encyclopedia Britannica,' 11th ed., vol. 7, p. 581, 1910; 14th ed., vol. 6, p. 819 (1929).

³⁰ Abich, H., Ann. d. Physik (Poga.), 146, 475-482 (1872).

⁸¹ Brewster, Sir David, Phil. Mag. (3), 4, 245-246 (1834).

⁸² Clarke, E. D., Trans. Cambr. Phil. Soc., 1, 209-215 (1822).

²⁸ Leydolt, F., Sitz. Akad. Wiss. Wien (Math.-nat.), 7, 477-487 (1851).

reports of hexagonal forms, such as those by W. A. Bentley, ¹⁶ T. H. Holland, ³⁴ P. A. Secchi, ³⁵ J. Smithson, ³⁶ and J. Tyndall, ³⁷ are probably to be interpreted as indicating that the crystals belonged to the hexagonal system, without indicating to which of the two divisions of that system they should be assigned.

Although Leydolt found ice crystals of only a single system and doubted the existence of other types, there are reports indicating that ice-I may, under conditions not yet defined, crystallize in the cubic system. For example, A. E. Nordenskjöld 38 has observed in the frost coating a windowpane, the outer air being at -8 to -12 °C, small rectangular forms which he decided could not possibly belong to the hexagonal system, but probably belonged to the rhombic, or possibly to the cubic. Similar forms have been observed by H. P. Barendrecht 39 to separate at low temperatures from solutions of water in acetaldehyde and in certain alcohols, including ethyl alcohol. By polariscopic observations, he found that these forms belonged to the cubic system. It was not practical to separate them from the viscous mother-liquor, but from the fact that the same type of crystal was obtained from all these solutions, he concluded that the crystals were ice and not a hydrate. Similar observations have been reported by F. Wallerant 40 who interprets them as indicating that the cubic crystals pertain to a type of ice that is stable only under high pressure. He remarked that in an alcoholic solution the cubic crystals are stable as long as the water content does not exceed 55% by weight; but if it does exceed that value, the cubic crystals, initially formed by supercooling the solution, transform into rhombohedric crystals of ordinary ice enclosing numerous (une infinité) small isotropic crystals. P. Tschirwinsky 41 has questioned the validity of the conclusions of Barendrecht and of Wallerant; he believed that the cubic crystals they obtained were those of the hydrates, such as $C_2H_5(OH) + 3H_2O$ of which the existence, he said, had been shown by Mendelejeff in 1865. It contains 54.1 per cent of water, which essentially coincides with the 55 per cent limit found by Wallerant for the stability of the crystals. R. Hartmann 42 also has reported a rectangular type of crystal skeleton obtained from aqueous solutions; but from the observation that the "freezing point" of supercooled water inoculated with such skeletons is the same as when it is inoculated with a crystal of the ordinary hexagonal type, he concluded that the skeletons actually belong to the hexagonal system.

⁸⁴ Holland, T. H., Nature, 39, 295 (1889).

³⁵ Secchi, P. A., Bull. meteor. Osserv. Coll. Romano, 15, 73-74 (1876).

⁸⁰ Smithson, J., Ann. Philos. (N. S.), 5, 340 (1823).

²⁷ Tyndall, J., "The Forms of Water in Clouds and Rivers, Icc and Glaciers," New York, D. Appleton & Co., 1872.

⁸⁸ Nordenskjöld, A. E., Ann. d. Physik (Pogg.), 114, 612-627 (1861).

³⁰ Barendrecht, H. P., Z. physik. Chem., 20, 234-241 (1896); Z. anorg. Chem., 11, 454-455 (1896).

⁴⁰ Wallerant, F., Bull. Soc. Franc., Minéral., 31, 217-218 (1908).

⁴¹ Tschirwinsky, P., Ann. Geol. et Min. Russie (French résumé), 14, 280-282 (1912) $\rightarrow N$. Jahrb. Min., Geol., Paläon., 19142, 349 (1914).

⁴⁹ Hartmann, R., Z. anorg. Chem., 88, 128-132 (1914).

The subject has been reviewed by O. Mügge ⁴⁸ and briefly summarized by W. H. Barnes ^{43a} who states that a very complete review, containing more than 100 citations, may be found in A. B. Dobrowolski's "Historja Naturalna Lodu."

J. M. Adams ⁴⁴ has stated that the ice-crystal is asymmetric with respect to its basal (0001) plane, crystals twinned on that plane being separable into two kinds: (a) Those that may develop pits at the ends of the c-axis, and (b) those that may develop a cavity at the middle of that axis. And J. Smithson ⁸⁶ has stated that when hail is sufficiently regular for satisfactory measurement, it always consists of two hexagonal pyramids joined base to base. "One of the pyramids is truncated," and "The two pyramids appeared to form by their junction an angle of about 80 degrees."

From his study of the way ice yields to stresses of various kinds, I. C. McConnel 45 found that "a crystal behaves as if it were built up of an infinite number of indefinitely thin sheets of paper fastened together with some viscous substance which allows them to slide over each other with considerable difficulty; the sheets are perfectly inextensible and perfectly flexible. Initially they are plane and perpendicular to the optic axis; and when by the sliding motion they become bent, the optic axis at any point is still normal to the sheet at that point." It will be noticed that these sheets are parallel to the planes of Tyndall's flowers of ice (p. 405), and to those in which he observed melting to occur when ice is subjected to linear compression along the optic axis (p. 431). Quincke stated that "the planes of easiest cleavage in natural ice crystals (laminated structure, displacement without bending) are due to invisible layers of liquid salt solution which are embedded in the crystals, normal to the optic axis, or often in other positions." 46 Whether the sheets of McConnel's are to be identified with certain of these layers of Quincke's is not clear.

Structure of Ice in Bulk.

However uniform the ice may be, melting begins at the boundaries between the individual ice crystals, and as it proceeds, the crystals become more and more separated one from another, and the ice becomes "rotten." There is between the crystals a material with a lower melting point than the crystals themselves, and this material, when molten, dissolves the surface of the ice-crystals in contact with it, even when its temperature is below the normal melting point of the crystal. This material surrounds each crystal, enclosing it in a cell. These correspond to the foam-cells that are postulated in the theory of solidification that was developed by G. Quincke in a series of papers published some 30 years ago, and that was applied by him to explain the formation of ice and of glacier grains.⁴⁷

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<sup>48</sup> Mügge, O., Centralbl. Min., Geol., Paläon., 1918, 137-141 (1918).

<sup>48</sup> Barnes, W. H., Proc. Roy. Soc. (London) (A), 125, 670-693 (1929).

<sup>44</sup> Adams, J. M., Proc. Roy. Soc. (London) (A), 128, 588-591 (1930) → Phys. Rev. (2), 36, 788 (A) (1930).
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McConnel, J. C., Proc. Roy. Soc. (London), 48, 259-260 (1890); 49, 323-343 (1891).
 Quincke, G., Paragraph 35 in Nature, 72, 543-545 (1905); Proc. Roy. Soc. (London) (A), 76, 431-439 (1905).

A brief outline of Quincke's hypothesis is given by H. T. Barnes.⁴⁸ On this hypothesis, the liquid contains at least two species of molecules, one vastly in excess of the other. The species present in smaller amount he regards as salt, but that is not at all necessary. Under the action of the intermolecular forces, there will, in general, be a segregation of each species; and those present in small amounts may form a net-work throughout the liquid, enclosing the more abundant species of molecules in numerous adjacent cells, resembling cells of foam. In the case of water, the contents of the cells (the purer water) freezes first; then the cell-walls freeze. Each individual crystal, each glacier grain, corresponds to a single cell. As the temperature rises, the cell-walls melt first. The resulting liquid (an aqueous solution) bathes the surfaces of the enclosed crystals, lowering their melting point, and so causes those surfaces to melt at a temperature below the normal melting point of the crystal itself. Thus, the crystals become separated more and more from one another.

Quincke has advanced the very extreme suggestion that "ice is a liquid ielly, with foam-walls of concentrated oily salt solution, which enclose foamcells containing viscous, doubly refracting, pure or nearly pure water. The further the temperature falls below 0°, the greater is the viscosity of both liquids—in the walls and in the interior of the foam-cells—and the less the plasticity of the ice. Ice crystals at temperatures below 0° consist of a doubly refracting viscous liquid, and are intermediate between the soft crystals of serum albumen and ordinary crystals of quartz, felspar, etc."49

Whether these cell walls (Quincke) generally exist in water before freezing begins may be open to question. But as soon as freezing begins they will certainly begin to form. When each small volume of water crystallizes, it rejects the impurities it originally contained. Each minute crystal is surrounded by a layer of water more impure than that from which it was formed. As the crystal grows, this layer is continuously pushed out, becoming ever more impure, but always hugging the crystal, until it meets a similar layer surrounding a neighboring crystal. Further growth in that direction means a thinning of the combined layer and an increase in its concentration. Thus each crystal becomes enclosed in a material having a lower melting-point than itself. Diffusion will, of course, tend to equalize at each instant the concentration of the impurities throughout the volume that is still liquid; and adsorption may tend to retain at the crystal surface more of one type of impurity than of another.

The actual existence of an intercrystallic material differing in properties from the crystals is conclusively shown by the fact that melting always begins at the boundaries between the crystals. Furthermore, E. K. Plyler 50

⁴⁷ Quincke, G., Ann. d. Physik (4), 18, 1-80 (1905); summary of conclusions is given in Proc. Roy. Soc. London (A), 76, 431-439 (1905) and in Nature, 72, 543-545 (1905).

⁴⁸ Barnes, H. T., "Ice Formation," p. 79-82.

⁴⁹ Quincke, G., Paragraphs 3, 4, 36 of Nature, 72, 543-545 (1905), and Proc. Roy. Soc. (London) (A), 76, 431-439 (1905).

⁵⁰ Plyler, E. K., J. Elisha Mitchell Soc., 41, 18 (1925).

observed that infrared radiation is much more strongly absorbed in the region between the crystals than in the crystals themselves; in some cases the boundary region was found to be less than 8μ (0.0008 cm) thick. cleaves most readily between crystals, that is, along the cell walls. The hypothesis of such cell walls accords with certain observation reported by G. Beilby,⁵¹ and with the observation that "the more slowly artificial ice is frozen, and the less salt it contains, the more transparent it is, and rigid, and the more difficult to split with a knife." 15, p. 19; 47, p. 79

The purer the water, the smaller is the amount of intercrystallic material, but that one is ever justified in considering that material as negligible seems improbable, especially when one realizes that the action of this material in separating the crystals is a surface phenomenon, and recalls how minute an amount of an impurity may suffice to produce a profound change in the properties of a surface.

Ouincke 47, pp. 78, 79 states that by repeated fractional freezing, with a discarding of the unfrozen fraction, the ice becomes increasingly purer and composed of ever larger grains, but he has never succeeded in obtaining ice without grains; i.e., with no intercrystallic material.

- J. N. Finlayson 52 writes: "The crystals first formed continually enlarge and the interstices between them become filled with smaller crystals not so regularly oriented. The crystals, however, do not completely unite and there is a definite cleavage plane formed between them."
- J. Y. Buchanan 53 has shown that the crystals of ice formed from a dilute salt solution are themselves free of salt, but that some of the solution is retained in the interstices between the crystals; at the same time he called attention to the pronounced effect this intercrystallic material may have upon the physical properties of the ice, even when the amount of salt present is excessively minute. The presence of such intercrystallic material should always be remembered when ice in bulk is being studied. In many cases it profoundly affects the results obtained.

The slow disintegration of ice when exposed to light and air near 0 °C has been described by E. Schmid.⁵⁴ It is frequently stated that such disintegration does not occur when the ice is completely submerged; but M. Faraday 55 observed that ice that was completely submerged in water contained in a vessel surrounded by an ice-jacket, and thus kept so near 0 °C that a cubic inch of ice was not dissolved in a week, became after several days "so dissected at the surfaces as to develop the mechanical composition of the masses, and to show that they were composed of parallel layers about a tenth of an inch thick, of greater and lesser fusibility, which layers appear, from other modes of examination, to have been horizontal in

⁵¹ Beilby, G., "Aggregation and flow of solids," pp. 140-143, 1921.

⁵² Finlayson, J. N., Canadian Engineer, 53, 101-103 (1927).

²⁵ Buchanan, J. Y., Proc. Roy. Soc. Edinburgh, 14, 129-149 (1887) → Nature, 35, 608-611 (1887); 36, 9-12 (1887); Proc. Roy. Inst. Grt. Brit., 19, 243-276 (1908).

⁵⁴ Schmid, E., Ann. d. Physik (Pogg.), 55, 472-476 (1842).

⁵⁵ Faraday, M., Proc. Roy. Soc. (London), 10, 440-450 (1860).

the ice whilst in the act of formation." However, L. Hawkes ⁵⁶ has stated that F. J. Hugi ⁵⁷ observed that "in cloudy weather a lump of ice melts as a whole, preserving a smooth outer surface, but in the sun's rays melting takes place at the intergranular boundaries." Such apparently incompatible observations are probably to be explained by the vast difference in the rates of melting.

A. Erman ⁵⁸ has reported many interesting and unusual observations made on ice in Siberia.

Internal Melting.

When a block of ice that is above water is exposed to light, small cavities partly filled with water may be seen to form throughout the body of the ice, provided that the temperature of the ice is not too low. The most noted and beautiful of these are Tyndall's "flowers of ice," which will be considered in the next section. At the same time, water will be formed in cavities that previously had been dry. As the melting proceeds, the crystals become separated one from another, and each crystal becomes split into laminas perpendicular to the optic axis.⁵⁹ In speaking of the splitting of glacier grains into such laminas, Buchanan says (p. 263): "It is only the grains that are exposed to the sky, and above water, that are so analysed; and prolonged exposure of this kind reduces a grain to the last stage of dilapidation. The grains beneath the surface, whether of ice or water, are almost completely unattacked." (Cf. Plyler's observations on the absorption of the infrared, p. 403.)

In speaking of melting in the interior of the blocks into which the ice becomes broken up, E. Schmid ⁵⁴ said that small bubbles form first, and then thread-like cavities grow out from them. Could he have been speaking of what Tyndall later called flowers of ice?

If a cavity contains an inclusion, such as soot, which is a good absorber of radiant energy, the melting is easily understood. The inclusion abstracts energy from the radiation, becomes heated, and melts the ice. The same explanation was proposed for the melting when the only inclusion is air; but J. Tyndall ⁶⁰ showed that the absorptivity of air is entirely too small for it to act in that way.

Nevertheless, internal melting occurs; and "proves that the interior portions of a mass of ice may be melted by radiant heat which has traversed other portions of the mass without melting them."

Tyndall suggested that the localization of internal melting at air-bubbles depends upon the existence of the free surface surrounding the bubble, that it is a surface phenomenon (see below); G. Quincke ⁴⁷ attributed all inter-

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66 Hawkes, L., Gcol. Mag., 67, 111-123 (1930).
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⁸⁷ Hugi, F. J., Edinb. New. Phil. J., 10, 337-338 (1831).

⁵⁸ Erman, A., Phil. Mag. (4), 17, 405-413 (1859).

⁵⁰ Buchanan, J. Y., Proc. Roy. Inst. Grt. Brit., 19, 243-276 (1908).

⁶⁰ Tyndall, J., Proc. Roy. Soc. London, 9, 76-80 (1858) → Ann. d. Physik (Pogg.), 103, 157-162 (1858).

nal melting to the presence of small amounts of impurities, which accords with Buchanan's conclusions ⁵³; and the Thomson brothers ⁶¹ were of the opinion that internal stresses were of prime importance. All may contribute, their relative importance varying with the conditions, but in very many cases the stresses are surely the most important. But whatever may be their practical value, Tyndall's views are interesting and suggestive. They should be compared with those of Faraday regarding regelation (p. 412).

From his carefully described observations on the gradual liquefaction of masses of ice by the formation and growth of drops of water within them, Tyndall inferred that the melting temperature differs from point to point, oscillating about its normal value. He wrote: "Through weakness of crystalline structure, or some other cause, some portions of a mass of ice melt at a temperature slightly under 32 °F., while others of stronger texture require a temperature slightly over 32° to liquefy them. The consequence is, that such a mass, raised to the temperature of 32°, will have some of its parts liquid and some solid." These variations in the melting point are attributed by W. Thomson (Lord Kelvin)^{61, p. 141+} solely to variations in the stress. That some portions melt below 0 °C is certain, and accords with the conclusions of Quincke 47, p. 17 and of Buchanan; but whether any portions not subjected to such a tension as exists at the boundaries of a cavity completely filled with the melt derived from the ice that originally filled it can remain solid above 0 °C is another question. Tyndall thought they could, provided they have no free surface.

This he endeavors to explain as follows: "Regarding heat as a mode of motion, the author [Tyndall] shows that the liberty of liquidity is attained by the molecules at the surface of a mass of ice before the molecules at the centre of the mass can attain this liberty. Within the mass each molecule is controlled in its motion by the surrounding molecules. But if a cavity exists at the interior, the molecules surrounding that cavity are in a condition similar to those at the surface; and they are liberated by an amount of motion which has been transmitted through the ice without prejudice to its solidity. The author proves, by actual experiment, that the interior portions of a mass of ice may be liquefied by an amount of heat which has been conducted through the exterior portions without melting them."

Flowers of Ice.

J. Tyndall 60 observed that when a beam of light is passed through a block of ice, its path rapidly becomes dotted with numerous points resembling shining bubbles of air. Their appearance is accompanied by a clicking sound. When examined with a lens, these "flowers of ice" are found to consist of a bright central spot surrounded by six coplanar petals composed of water. Neighboring flowers lie in parallel planes which are independent

⁶¹ Thomson, J., Proc. Roy. Soc. (London), 8, 455-458 (1857); 10, 152-160 (1859); 11, 198-204 (1861); 11, 473-481 (1861); Thomson, W. (Lord Kelvin), Idem, 9, 141-143 (1858); 9, 209-213 (1858).

of the direction of the beam of light. It has been found that the plane of each flower is perpendicular to the optic axis of its associated crystal, and that the petals are parallel to the secondary axes of the crystal. Hence, they serve as a simple means for determining the orientations of the several individual crystals. Tyndall stated that the flowers are formed in planes parallel to those of freezing, but that some apparent exceptions had been noted. He goes on to say: "In some masses of ice, apparently homogeneous, the flowers were formed on the track of the beam, in planes which were in some cases a quarter of an inch apart."

- E. Hagenbach-Bischoff, sometimes referred to as Hagenbach, stated that the flowers are sometimes circular, and that they ordinarily begin as circles, that the plane of a flower is perpendicular to the optic axis, and that its arms are parallel to the secondary axes of the crystal.⁶²
- G. Quincke ⁶³ has stated that "at the edge of Tyndall's liquefaction figures, while they are in the process of enlarging, or on the bursting of the foam-walls of artificial ice as it melts, one often sees periodic vortex movements. These arise from a periodic capillary spreading out (Ausbreitung) of the salt solution of the foam-walls at the boundary between pure water and air or vacuum." It seems probable that those movements are very closely related to those observed by Tyndall when he compressed ice in the direction of the optic axis (see p. 431).

The clicking that accompanies the formation of the flowers was explained by Tyndall somewhat as follows: As the melting progresses the water formed adheres to the walls and so is subjected to tension, its volume being normally less than that of the ice from which it is formed. As soon as rupture occurs at any point, the tension is at once relieved, and the water immediately contracts, leaving a vacuous cavity. This sudden contraction gives rise to the click that is heard.

- J. M. Adams and W. Lewis 66 have observed that when the water in such a cavity is refrozen it "never completely fills the space, but at the center of each pattern, there remains a group of minute cavities precisely bounded by natural faces." These they call "negative crystals." "The question as to the source of the space occupied by the negative crystals remains for the present unanswered." These remarks refer more particularly to cavities that have grown beyond the point at which the flowers of ice are well formed.
- L. Hawkes ⁵⁶ has suggested that the flowers of ice mark the positions of particles of dust included in the ice, the melting resulting from the radiation absorbed by the dust.

For additional information see note. 65

Hagenbach-Bischoff, E., Arch. des Sci. Phys. et Nat., Genève (3), 23, 373-390 (1890).
 Quincke, G., Paragraph 37 of Nature, 72, 543-545 (1905); Proc. Roy. Soc. (London) (A), 76, 431-439 (1905).

[©] Hess, H., "Die Gletscher," p. 13, 1904; Barnes, H. T., "Ice Engineering," p. 18, 19; "Ice Formation," p. 77, 78; Trans. Roy. Soc. Canada III (3), 3, 3-27 (22) (1909); Hagenbach-Bischoff, E.

⁶⁶ Adams, J. M., and Lewis, W., Rev. Sci. Inst. (N. S.), 5, 400-402 (1934).

Formation of Frazil, or Needle Ice.

The fine needle-like ice that is often distributed throughout the volume of rapidly moving water is frequently called "frazil." It causes much trouble in the operation of hydraulic power plants. Its formation has been described in some detail by H. T. Barnes 67 in terms of his colloidal theory. 68 He states that when the growing ice particles can be first seen microscopically they are disk-like, and devoid of crystal form; they flocculate and grow into true crystals. In rapidly moving water the crystals are broken up before they can become large, and are thoroughly mixed with the water. giving it a cloudy appearance. This is the beginning of frazil. It forms throughout the body of the river if the water is supercooled, even if the supercooling does not exceed a few thousandths of a degree. It is adhesive, forms agglomerations, and is carried under the surface ice, forming hanging dams, which may reach down to the river bed. When the temperature of the water is a few thousandths of a degree below 0 °C the frazil aggregates are very strong and tenacious, but when the water is at or above 0 °C they become soft and spongelike. He states that sunlight not only warms the water, but has a direct action on the colloidal ice particles, destroying their "agglomerating properties." It seems probable that the adhesion and agglomeration of the particles of frazil are related to the phenomena described by L. Dufour. 69 For additional information regarding colloidal and frazil ice, see the articles already cited and H. T. Barnes 70 and P. P. von Weimarn and W. Ostwald.71

Formation of an Ice Sheet.

(For the freezing of water cooled much below 0 °C, see p. 416.)

The formation of surface ice has been described in detail by H. T. Barnes, 15, pp. 60, 61 H. Hess, 65, p. 11 G. Tammann and K. L. Dreyer, 72 G. Seligman, 73 and others.

As the water of a pond is being cooled it is warmer than the air; consequently currents of warm air rise from the center of the surface, and cold air sweeps in from the sides. This chills the lateral waters, and is itself warmed thereby; thus the center of the surface remains warmer than the edge, and if the air temperature is not very low, the center may remain unfrozen long after the banks are bordered with ice. Until the temperature of the water has dropped to 4 °C, convection keeps it fairly uniform throughout the depth; with further chilling, the colder layers remain at the top. If surface freezing is to occur, either the temperature of the air

⁶⁷ Barnes, H. T., "Ice Engineering," p. 6+, 108, 109; Scientific Monthly, 29, 289-297 (1929). See Barnes, H. T., "Colloid Chemistry," J. Alexander, ed., vol. 1, pp. 435-443, New York, Reinhold Publishing Corp., 1926.

Dufour, L., Ann. d. Physik (Pogg.), 114, 530-554 (1861).

⁷⁰ Barnes, H. T., "Colloid Symposium Monograph," Vol. 3, 103-111, New York, Reinhold Publishing Corp., 1925; "Ice Formation," 1906.

⁷¹ von Weimarn, P. P., and Ostwald, W., Koll. Z., 6, 181-192 (1910).

⁷² Tammann, G., and Dreyer, K. L., Naturwissenschaften, 22, 613-614 (1934).
72 Seligman, G., Proc. Roy. Inst. Grt. Brit., 29, 463-483 (1937) → Nature, 139, 1090-1094 (1937).

must be well below 0 °C, or the loss of heat by direct radiation must be great; otherwise the surface will receive heat from the lower layers more rapidly that it can lose it. When the temperature of the surface water has dropped sufficiently (Barnes says to 0 °C, but it probably drops appreciably below that 72; crystallization begins at the banks, and needles of ice shoot out over the surface of the water. These branch and broaden until the entire surface of the water along the banks is covered with a thin layer of ice. How far this layer extends toward the center depends upon the existing conditions. If the entire surface is supercooled at the time the crystallization begins, the needles may extend rapidly throughout the entire surface, which will soon be completely covered with ice. Under other conditions the initial layer of ice will not reach the center of the surface, but will gradually grow toward it. When the surface has become iced over, the ice layer gradually thickens as heat passes out by conduction through it. H. T. Barnes 74 has stated that the growth in thickness is by an accumulation on the underside of the sheet of layers of disks of ice, "very much like stacked Chinese coins." Some have thought the initiation of freezing at the bank, instead of elsewhere, arises from an increased cooling caused by conduction of heat from the water and through the shore material, but in view of the low conductivity of rock, sand, and clay, it seems that this effect must be exceedingly small as compared with that due to the heat carried away convectively by the air.

These descriptions must be supplemented by important observations made many years ago by F. Klocke. 75 He observed that, if the chilling is severe, the lengths of the needles that shoot out over the surface as the water begins to freeze are parallel to their optic axes; for these needles the optic axis is parallel to the surface of the water. The needles broaden, on one side mainly; this is not an outgrowth of the needle. The optic axis of this newer ice is perpendicular to the surface. The main sheet of ice from the beginning consists of this newer ice, the needles seeming to be extraneous impurities. If the freezing occurs near 0 °C, the optic axes of even the needles are perpendicular to the surface.

A. E. Nordenskjöld ⁷⁶ reports that, on August 31, 1878, when the sky was clear except near the western horizon, and the temperature of the water near the surface was between +1 and +1.6 °C, and that of the air on the vessel between +1.5 and +1.8 °C, "ice was seen to form on the calm, mirror-bright surface of the sea. This ice consisted partly of needles, partly of a thin sheet. The formation of (this) ice was clearly a sort of hoar-frost phenomenon, caused by radiation of heat."

After the surface of quiet water has become covered with ice, the sheet has been found to grow in thickness (x) in accordance with the empirical

⁷⁴ Barnes, H. T., Scientific Monthly, 29, 289-297 (1929).

Klocke, F., Jahrb. Mineral., Geol., 1879, 272-285 (1879).
 Nordenskjöld, A. E., "The Voyage of the Vega around Asia and Europe," New York, Macmillan & Co., pp. 317, 318, 1882.

formula $x + x^2/2 = -\frac{\tau Kt}{LS}$, where τ is the time, t the temperature of the

air, K the thermal conductivity of ice = 0.0057 cal/cm²-sec per °C/cm, L the latent heat = 80 cal/g, and S the density of ice = 0.9166 g/cm³.¹⁵, p. 63 Whence we get Table 188. This formula presumably supersedes the one, $x^2 = -K\tau t$, given by P. Vedel,⁷⁷ who recommended an experimental determination of K in each particular case.

Table 188.—Rate of Thickening of Ice-Sheet

If the temperature of the air is t °C and that of the water is 0 °C, τ is the time required for the sheet to become x cm thick. The sheet is assumed to be free from snow. (See text.)

2 2.9 hr 1.4 hr 43 min 29 min 21 10 1.79 da 21.4 hr 10.7 hr 7.1 hr 5 15 3.80 da 1.90 da 22.8 hr 15.2 hr 11. 20 6.55 da 3.28 da 1.64 da 26.2 hr 19. 30 14.29 da 7.15 da 3.57 da 2.38 da 1.	-40	
	8.0 min 1 min 5.4 hr 1.4 hr 9.7 hr 1.79 da 6.92 da 5.4 da	

Growth and Orientation of Crystals.

U. Yoshida and S. Tsuboi ⁷⁸ report (a) that all directions parallel to the basal plane of the hexagonal crystal of ice are equally suited for growth, which occurs more readily in these directions than in a direction that is perpendicular to that basal plane;* (b) that ice formed on the surface of calm water exposed to cold air during a fine night is usually composed of monocrystals of considerable size with their basal planes nearly parallel to the surface; (c) that ice columns formed in the ground usually take the form of long prisms, each consisting of several smaller prisms of monocrystals about 0.5 mm. in diameter (cf. p. 419); and (d) that the lower end of an icicle usually consists of a slender monocrystal. In all these cases the growth is parallel to the basal plane of the crystal. If a monocrystal of ice only slightly below 0 °C is placed in contact with a drop of water, the water freezes slowly, and the crystallographic axes of the new ice are "entirely the same" as that of the mother crystal (cf. p. 411).

Likewise, F. Leydolt ³³ reported, as a conclusion from his extended polariscopic study of ice, that the optic axis of the ice is always perpendicular to the surface of the ice-sheet, whether that has been formed on a river, a pond, or in a small vessel. And still earlier, G. Tammann and

^{*} H. D. Megaw * has stated that the direction of fastest growth is along the normal to the plane (11 $\bar{2}0$).

⁷⁷ Vedel, P., J. Franklin Inst., 140, 355-370, 437-455 (1895).

⁷⁸ Yoshida, U., and Tsuboi, S., Mem. coll. sci., Kyoto (A), 12, 203-207 (1929).

⁷⁹ Megaw, H. D., Nature, 134, 900, 901 (L) (1934).

K. L. Dreyer,⁷² and G. Seligman ⁷³ had reported likewise; and R. Mallet ⁸⁰ had announced that if any crystallizable material is "suddenly cooled from a state of fusion or solution, by a plane surface of low temperature, the crystals in forming arrange themselves perpendicularly to the refrigerating plane," in the direction of the flow of heat.

The orientation of the crystals in ice formed on the surface of calm water is generally reported to be as just described.⁸¹ But I. C. McConnel and D. A. Kidd 81a have observed surface ice in which the crystals were arranged with the optic axis nearly horizontal, and hence with the basal planes nearly perpendicular to the surface. They write (p. 334): "Some of the ice of the St. Moritz lake is built up of vertical columns, from a centimeter downward in diameter, and in length equal to the thickness of the clear ice, i.e., a foot or more. A horizontal section, exposed to the sun for a few minutes, shows the irregular mosaic pattern of the divisions between the columns. The thickness of each column is not perfectly uniform. Sometimes indeed one thins out to a sharp point at the lower end. Each column is a single crystal, and the optic axes are generally nearly horizontal. Some experiments on freezing water in a bath lead us to attribute this curious structure to the first layer of ice having been formed rapidly, in air, for instance, below -6 °C. We found that if the first layer had been formed slowly, and was therefore homogeneous with the axis vertical, a very cold night would only increase the thickness of the ice while maintaining its regularity."

In the usual case of an ice sheet formed on calm water, the optic axes of the crystals are both perpendicular to the refrigerating surface and parallel to the direction of gravity. Whence have arisen two explanations of this orientation. A. Bertin 82 announced that the direction of the optic axis is determined by that of the refrigerating surface, being always perpendicular to that; and in confirmation he reported certain observations in which the refrigerating surface was inclined to the horizontal. His conclusions have been confirmed by F. Klocke. To On the other hand, O. Mügge 81 maintained that the direction was determined gravitationally in the case of calm water; the most rapid growth taking place in the basal plane, plates normal to the optic axes are formed, and these will, obviously, float with the optic axes vertical. If there are currents in the water, as will be the case if the refrigerating surface is not horizontal, these currents will affect the orientation of the plates. In such a way he attempts to set aside Bertin's observations.

It seems most probable that Bertin's view is correct, that as the ice crystal is initially formed its optic axis is perpendicular to the refrigerating surface—or more specifically, it is parallel to the temperature gradient—

⁸⁰ Mallet, R., Phil. Mag. (3), 26, 586-593 (1845).

⁸¹ See also, Mügge, O., Newes Jahrb. Mineral., Gcol., 1895_{II}, 211-228 (1895); Reusch, E., Ann. d. Physik (Pogg.), 121, 573-578 (1864); Bertin, A., Ann. de chim. et phys. (3), 69, 87-96 (1863); von Engeln, O. D., Am. J. Sci. (4), 40, 449-473 (1915); Finlayson, J. N.⁶²

⁸¹a McConnel, J. C., and Kidd, D. A., Proc. Roy. Soc. (London), 44, 331-367 (1888).

⁸² Bertin, A., Ann. de chim. et de phys. (5), 13, 283-288 (1878).

except in so far as mechanical or other disturbances cause it to take some other direction. When the crystal is formed on the surface of calm water, the gravitational effect mentioned by Mügge will cooperate in keeping the axis vertical; currents in the water may force the axis out of line with the temperature gradient. When crystallization begins on the surface of water that has been cooled significantly below 0 °C, it will give rise to temperature gradients in the plane of the surface, and these may account for the fact that the optic axes of the needles that first shoot out over the surface are horizontal, as noted by F. Klocke (p. 408), and that the crystalline structure of the first layer of ice is irregular (Bertin). Whether the same explanation is applicable to the ice, with optic axis horizontal, observed by McConnel and Kidd (p. 410) is not clear.

These unqualified conclusions apply only to the initial crystals formed on the surface of water previously free from ice. If there are ice crystals already present—blocks of ice, snow crystals—then a new effect comes into play: namely, the tendency for the new crystal to take the orientation of the old crystal from which it springs. The actual orientation will depend upon the relative magnitudes of the two tendencies and upon the difference in the orientations to which they would individually give rise. J. Y. Buchanan bas described the irregular crystalline structure of surface ice initiated by a floating block of glacier-ice.

In the preceding paragraphs we have considered the direction of the optic axis only, and have seen that under certain conditions, frequently realized, the individual crystals over a considerable expanse of ice are so oriented that their optic axes are parallel, or very nearly so. Turning to the secondary axes we find no such regularity (Mügge 81). The orientation of neighboring flowers of ice, which indicate the orientation of the secondary axes, may be most varied.

K. R. Koch 84 has suggested that it is very probable that the orientation of the individual crystals may be uniform only in thin layers of ice, and that, in a thick block, it may differ from layer to layer. This seems to refer to the optic axis as well as to the secondary axes.

U. Yoshida and S. Tsuboi ⁷⁸ state that when the freezing is very rapid, the ice crystals are oriented at random, even when the freezing is induced

by a mother crystal.

When frost forms on ice, the optic axis of each little crystal added by the vapor is parallel to that of the ice crystal to which it is attached.⁸⁵ In general, the optic axes of adjacent crystals of ice are not parallel, in which case, the frost crystals point in various directions. The ice crystals formed by condensation on highly chilled metal plates are oriented at random (Yoshida and Tsuboi ⁷⁸).

C. R. Elford 86 has published photographs of triangular crystals extend-

See also, Barnes, H. T., Nature, 83, 276 (1910); von Engeln, O. D., Am. J. Sci. (4), 40, 449-473 (1915).

⁸⁴ Koch, K. R, Ann. d. Physik (4), 45, 237-258 (1914). 88 Plyler, E. K., J. Opt. Soc. Amer.. 9, 545-555 (1924).

Plyler, E. K., J. Opt. Soc. Amer., 9, 545-555 (1924).
 Elford, C. R., Monthly Weather Rev., 64, 83 (1936).

ing downward from heavy surface ribs in the ice sheet that formed on a mud-puddle that had dried up during the night.

Recrystallization.

If a vitreous solid is maintained above a certain temperature, generally well below its melting point, it will in time become crystalline. This limiting temperature, below which the change will not occur, is called the temperature of recrystallization. When a differential stress causes the substance to flow while at a temperature between that of recrystallization and the melting-point, the flowed substance recrystallizes as soon as the stress is relieved; but if the temperature is below that of recrystallization, the flowed material assumes the vitreous state. The temperature of recrystallization of ice is considerably below $-12\,^{\circ}\text{C.}^{87}$ For an example of marked recrystallization caused by stress, see O. D. von Engeln. 81 See also p. 438.

R. Mallet ⁸⁰ has written: "If a crystallizable body be heated near to but not up to its fusing point, by the application of heat in one plane, a crystalline structure perpendicular to the plane is immediately developed." And again: "In general, change of temperature beyond certain limits develops in crystallizable bodies a crystalline structure in the direction of the wave of heat, whether into or out of the mass of the body."

The term recrystallization is also used to denote the growth of one crystal at the expense of another. In order to distinguish this phenomenon from the preceding, it will be called migratory recrystallization.

G. Beilby ⁸⁸ failed to observe any indication of migratory recrystallization during the slow warming and melting of a film of ice initially at -11 °C, but G. Tammann and K. I. Dreyer ⁸⁹ seem to have observed migratory recrystallization of ice, and the fact that glacier grains increase in size and decrease in number as the ice ages and moves under complex stresses shows that, however it may arise, there is in that case such an effect. ^{62, 90} And H. T. Barnes ⁷⁴ has stated that surface ice becomes coarser with age, the large crystals consuming the smaller ones. Furthermore, J. Thomson ⁹¹ has shown that there must be migratory recrystallization whenever it is possible for molecules to pass by any means from a stressed crystal to one that is less stressed. An obvious means by which they may so pass is by fusion, or sublimation, and resolidification, the temperature being suitable.

Regelation.

M. Faraday,⁹² observed that when two pieces of ice, each at 0 °C, are brought into contact they freeze together, and do this even in a vacuum or

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87 Beilby, G., "Aggregation and Flow of Solids," p. 196, 1921.
88 Beilby, G., "Aggregation and Flow of Solids," 1921.
89 Tammann, G., and Dreyer, K. I., Z. anorg. allgem. Chem., 182, 289 313 (1929).
90 Vallot, J., Compt. rend., 156, 1575-1578 (1913).
91 Thomson, J., Proc. Roy. Soc. (London), 11, 473-481 (1861).
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Faraday, M., Proc. Roy. Inst. Grt. Brit., 1850; Exp. Res. in Chem. and Phys., pp. 372-374, 377-382 (1859); Proc. Roy. Soc. (London), 10, 440-450 (1860).

in water, and when brought together with the least possible pressure. Tyndall called this "regelation." Both he and Faraday explained it on the hypothesis that a thin layer of water bounded on each side by ice will freeze when it would not do so under other conditions. Having set forth his hypothesis that superficial portions of ice melt at a lower temperature than do others, Tyndall 60 writes: "The converse of this takes place when two pieces of ice at 32 °F., with moist surfaces, are brought into contact. Superficial portions are by this act virtually transferred to the center; and as equilibrium soon sets in between the motion of the tenuous film of moisture between the pieces of ice and the solid on each side of it, the consequence is shown to be that the film freezes, and cements the two pieces of ice together."

This explanation was vigorously upheld by Faraday 92 and as vigorously attacked by the Thomson brothers,93 who ascribed regelation to a melting produced by stress, followed by solidification when the stress is relieved, such melting and refreezing having been shown by J. Thomson 91, 94 to be necessary consequences of varying stress. Even in a vacuum the surfaces are moist, and there is a pressure due to capillary action; when the ice is in water, there are pressures arising from the unavoidable currents set up when the two pieces are brought together; and in all cases there are pressures arising from mechanical disturbances transmitted from without. This explanation—melting produced by stress and followed by freezing when the stress is relieved—is the one now generally accepted.

In contrast to the preceding explanations, L. Pfaundler 95 thought that regelation was to be explained by the presence in melting ice of molecules of liquid as well as of solid, the two species continually changing one into the other, but just how he expected this to bring about the observed effect is not clear.

E. W. Brayley 96 thought that there was a close analogy between regelation and the union of polished plates of glass in the manufacture of plate glass. And that suggests a possible similarity between it and the adhesion between flat plates, whether of glass or of metal, that have been wrung together.

The completeness with which two blocks of ice will freeze together increases with the pressure and its duration, and depends upon the relative orientations of the crystals of the two blocks. If the orientations are exactly the same in the two blocks, then they freeze together completely. the plane of union differing in no respect from any other parallel plane in either block. If the principal axes are parallel, but the subordinate axes of the crystals in one block are not parallel to those of the crystals in the other. the blocks freeze together so completely that the plane of separation cannot

[∞] Thomson, J., Proc. Roy. Soc. (London), 10, 152-160 (1859); 11, 198-204 (1861); Thomson, W. (Lord Kelvin), Idem, 9, 141-143 (1858).

[∞] Thomson, J., Trans. Roy. Soc. Edinburgh, 16, 575-580 (1849).

^{*} Pfaundler, L., Müller-Pouillet's "Lehrbuch der Physik," 9th ed., vol. 22, p. 595, 1898; Cf. Sits. Akad. Wiss. Wien, 592, 201-206 (1869). 96 Brayley, E. W., Proc. Roy. Soc. (London), 10, 450-460 (1860).

be detected by polariscopic observations, but can be by the production of Tyndall's flowers of ice, which show the difference in the orientations of the crystals in the two blocks. When tested by compression, the block yields first in that plane if the temperature of the room is over 0 °C, but the initial yield bears no relation to that plane if the temperature is below 0 °C. If the principal axes of the crystals of one block are perpendicular to those of the other, the union may again be invisible, but its strength is less than in the preceding case.97

Purity of Ice.

- H. T. Barnes 98 has frequently emphasized the purity of ice, especially of that formed on the underside of a thick sheet growing over flowing water. That the great bulk of the impurities carried by the water is eliminated under these conditions seems certain, but the elimination of "every trace of foreign matter" would seem to require very special conditions.
- G. Quincke 47 stated that the purity of the resulting ice is continuously increased by successive fractionings by freezing; but that he never succeeded in obtaining ice that was free of grains, that formed a single crystal.
- F. Witt 99 has found that the proportion of radon included in the ice that first forms when water containing radon is frozen depends upon the rapidity of the freezing. When the freezing was as slow as 0.0005 cm³/sec, the concentration of the radon in the ice was only 3 or 4 per cent of that in the water; but if the rate of freezing exceeded 0.001 cm³/sec the amount increased rapidly with the rate.

Production of Homogeneous Ice.

In 1845, C. Brunner 100 described his attempts to produce homogeneous ice suitable for use in a determination of the density of ice. On cooling while exposed to the air, carefully boiled-out distilled water takes up so much air that when it is frozen it contains many bubbles, especially in the portion that was the last to freeze. Covering the water with turpentine immediately after the boiling kept out the air fairly well, but the ice was then full of cracks. Following a suggestion, which he attributes to F. C. Achard, 101 he exposed one side of the vessel, containing the water to be frozen, to a low temperature and the opposite to a temperature above 0 °C. The ice so formed contained some air-bubbles, but not nearly so many as that formed in the usual manner. He found that selected river-ice was much superior to any ice he succeeded in freezing in the laboratory.

L. Dufour 102 boiled-out and froze water in a Torricellian vacuum, air

⁹⁷ See Hess, H.⁶⁵, pp. 25, 26; Heim, A., Ann. d. Physik (Poyg.) Erg. Bd., 5, 30-63 (1871); Hagenbach-Bischoff, E.⁶²

Barnes, H. T., "Colloid Symposium Monographs," 3, 103-111, Reinhold Publishing Corp., 1925;
 "Colloid Chemistry" (J. Alexander, ed.), Vol. 1, pp. 435-443, 1926.
 Witt, F., Sitz. Akad. Wiss. Wien (Abt. IIa), 139, 195-202 (1930).

¹⁰⁰ Brunner, C., Ann. d. Physik (Pogg.), 64, 113-124 (1845).

¹⁰¹ Achard, F. C., "Chem.-Phys. Schriften," Berlin, 1780.

¹⁰² Dufour, L., Compt. rend., 54, 1079-1082 (1862).

pressure not over 0.5 mm. The ice contained a few very small bubbles. It was opalescent and very homogeneous. He states that the opalescence was not due to air, but to the crystalline structure of the ice, or to internal crevices.

- R. Bunsen 103 introduced boiled-out distilled water into one arm of a U-tube initially filled with air-free mercury. The water was introduced while hot, was boiled in the tube, and the tube was then sealed so as to exclude all air. The water was frozen gradually from the top downward. Thus he obtained a cylinder of ice that was entirely (völlig) free of air-bubbles, and that was equal to the best crystal glass in clearness and transparency.
- G. Forbes ¹⁰⁴ used the same procedure, and stated: "The ice formed was quite uniform, very clear, and when cloven by planes perpendicular to the plane of freezing, split easily, showing the crystalline structure with great clearness."
- G. Quincke ⁴⁷ stated that by repeated fractionations, by means of freezing and thawing, the ice becomes ever purer and purer, and the crystals larger, but that he had never succeeded in obtaining ice that was not an aggregate of many crystals. The removal of air by alternate freezing and rapid thawing had been described, and the method seems to have been used by Duvernoy ¹⁰⁵ nearly 40 years before.

A. Leduc ¹⁰⁶ introduced hot, boiled-out distilled water into an exhausted vessel, and froze it progressively from the bottom to the top. The upper portions contained bubbles. Even after three such freezings *in vacuo* there were small bubbles in the portion last frozen. A fourth freezing appeared to produce no further improvement. Likewise, G. Bode ¹⁰⁷ has stated that neither the boiling-out of distilled water nor its repeated freezing in a vacuum is sufficient to insure a clear sheet of ice.

H. Hess ¹⁰⁸ stated that by slow, long-continued freezing very homogeneous ice-sheets may be formed on the water in large reservoirs. On thawing, these sheets break up into vertical, columnar pieces.

Monocrystals of Ice.

A portion of ice is defined by J. M. Adams and W. Lewis ¹⁰⁹ as monocrystallic if the flowers of ice formed in it all lie in parallel planes and are similarly oriented. They actually use the faces of the negative crystals (p. 406), formed when the water in those cavities is refrozen, as indices of the orientation of the flowers.

They have reported that monocrystals "with dimensions of the order of 10 cm were readily produced" by the following method. A mono-

Bunsen, R., Ann. d. Physik (Pogg.), 141, 1-31 (1870).
 Forbes, G., Proc. Roy. Soc. Edinburgh, 8, 62-69 (1873).

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100 Duvernoy, Ann. d. Physik (Poga.), 117, 454-463 (1862).
100 Leduc, A., Compt. rend., 142, 149-151 (1906).
107 Bode, G., Ann. d. Physik (4), 30, 326-336 (1909).
108 Hess, H., "Die Gletscher," 1904.
100 Adams, J. M., and Lewis, W., Rev. Sci. Inst. (N. S.), 5, 400-402 (1934) → Phys. Rev. (2), 46, 328 (A) (1934).
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crystallic fragment of commercial ice is cut with two faces parallel and presenting any desired aspect to the principal axis. One of these faces is frozen to the outer surface of a metal vessel containing a freezing solution (-10 °C), and the opposite face is dipped just below the surface of distilled water at 0 °C in an ice-jacketed vessel. They boiled the water and cooled it rapidly to 0 °C. The main body of the growth is monocrystallic with the seed. As the length increases, growth becomes slow, and it may be desirable to make a new start, using a seed of greater cross-section, cut from the recently formed monocrystal. Parasitic crystals tend to start where the air-water surface meets the seed, and should be removed from time to time.

H. D. Megaw 79 has grown single crystals in a capillary tube of Lindemann glass in a copper-wire holder cooled by means of a mixture of acetone and solid carbon dioxide. And J. Meyer and W. Pfaff 110 have observed spicules of ice which they thought were monocrystals.

Freezing of Supercooled Water. (See also Section 97.)

That water can be cooled much below 0 °C without freezing, is well known, G. Oltramare 111 has stated that both R. Pictet and L. Dufour had carried the supercooling to -40 °C (no citation). The greater the supercooling the more readily does freezing occur, at least within certain limits, but the only certain way to initiate freezing intentionally is to "seed" the water with a suitable crystal. The smallest particle of ice will at once grow rapidly when placed in supercooled water, branching and spreading until either all the water has become frozen or the temperature has risen to 0 °C. All other methods commonly quoted as efficacious are so irregular in their actions, or have been shown to fail so signally under certain conditions, often ill-defined, that one must conclude that they are, at best, only secondarily involved in the initiation of the freezing that is sometimes, or often, observed when they are employed. On the other hand, certain conditions seem to oppose freezing. These are considered in Section 97. Initiation of freezing is not the mere negative of opposition to freezing, although the contrary might be inferred from many of the articles on this subject.

The freezing of water that has been greatly supercooled has been described by H. Hess, 108, p. 12 who states that the resulting ice generally takes the form of hexagonal prisms or plates, reminiscent of snow-crystals. H. T. Barnes and H. L. Cooke 112 had more difficulty in cooling water through 0 °C than in continuing the cooling to lower temperatures, and stated that when freezing occurred at low temperatures (approximately -10 °C) "the ice formed all through the mass of the water." They also reported that "care had to be taken that the thermometer bulb never touched the bottom of the flask when the water was supercooled, as it almost

Meyer, J., and Pfaff, W., Z. anorg. allgem. Chem., 224, 305-314 (1935).
 Oltramare, G., Arch. des sci. phys. et nat. (3), 1, 487-501 (1879).
 Barnes, H. T., and Cooke, H. L., Phys. Rev., 15, 65-72 (1902).

invariably caused freezing to take place. The ice would start at the point of contact and immediately spread out all through the mass of the water."

Cooling water to -10 or -13 °C is not so difficult as one might infer from the published accounts (see p. 638).

When water contained in a wide tube, or other vessel, is frozen rapidly, it is not unusual to see spicules of ice, from a millimeter to several centimeters in length, growing from the surface, or from the walls of the vessel. These occur when the surface of the water, including the film on the walls, is frozen first. This traps the remaining water, so that its pressure rises as further freezing occurs, until presently it breaks through at some point of weakness, gushing out in a jet that freezes at once into a tube that continues to grow at its tip. The break usually occurs in the surface ice, but the pressure may be transmitted up the film on the surface of the vessel to a weak place in the covering ice. 72, 118 If heat is being abstracted from the water rapidly, but only a small portion of the water was supercooled when freezing began, the growth of the spicule may be slow. But if at the initiation of freezing the bulk of the water was considerably supercooled, the spicule shoots out with a surprising velocity. Such growths have been observed by the compiler, and always under conditions that seem to demand the explanation just offered. On the other hand, J. Meyer and W. Pfaff 110 have described similar spicules which they think arise from condensation of the vapor, and which they believe are monocrystals.

Icicles.

No record of any careful study of the crystalline structure of icicles has come to the compiler's attention except that of F. Leydolt.³³ As a result of his polariscopic study of icicles, he announced that the optic axis is always radial, normal to the geometrical axis of the icicle. Nevertheless, other quite positive though mutually contradictory statements occur. It will suffice to mention four.

If the icicle is produced by slow and continuous freeing, it consists of a single crystal with its optic axis horizontal; and if it hangs from the eaves of a roof, its optic axis is not only horizontal, but is also perpendicular to the eaves. If the growth is interrupted by a drop in the temperature, and is later renewed by the thawing of ice or snow on the roof, then the icicle acts as the chilling surface, and the axes of the new crystals are everywhere perpendicular to the surface of the original icicle. So says H. Hess, 108, p. 12 but it is possible that his statements are colored by his belief that the optic axis is always parallel to the heat-stream at the instant the crystal is formed. The truth of the second sentence of this paragraph is especially doubtful as it seems more reasonable to expect that the axis of the new crystal will be parallel to that of the crystal on which it is being formed.

In marked contrast to the preceding, J. C. McConnel and D. A.

¹¹⁸ Bally, O., Helv. Chim. Acta, 18, 475-476 (1935).

Kidd 81a, p. 384 have stated: "An icicle is an example of ice formed of very minute crystals irregularly arranged." A. Bertin 82 said that the crystallization of an icicle is confused, and U. Yoshida and S. Tsuboi 78 have stated that the lower end of an icicle usually consists of a slender single crystal.

Hail.

Treatises and journals devoted to meteorology, as well as those devoted to physics, should be consulted by one interested in the nature of hail and in the conditions under which it is formed. The experiments and discussions by L. Dufour, 69, 114 G. Oltramare, 111 and K. C. Berz, 115 are interesting and suggestive. Some of Dufour's experiments are considered in Section 97.

Hailstones may be very large. G. Oltramare 111 has stated that they may weigh as much as 500 grams (no citation). Captain Blakiston 116 has reported an ice storm in which large stones fell. He weighed blocks of 3.5 and 5 ounces (100 to 140 g), and pieces the size of a brick were said to have been seen. P. A. Secchi 117 has described a violent hailstorm in which many of the hailstones consisted of clusters of hexagonal prisms terminated at their outer ends by pyramids [illustrations], some of the crystals being a centimeter long and correspondingly wide. Many of the clusters were 5 to 6 cm in diameter, and some weighed as much as 300 grams.*

The velocity of the uprush of air that is required to support a spherical hailstone of density 0.7 g/cm³ at a height of about 5 km above sea-level has been computed by W. J. Humphreys 118 to be as follows:

Diameter	1	2	2.5	3	3.5	4	5	inches
Velocity	55	78	91	109	136	185	219	miles/hour

He gives corresponding values for other densities between 0.9 and 0.5 g/cm³. Similar data have been given by M. A. Giblett ¹¹⁹ and by G. Grimminger.120

- J. Smithson 121 observed that hail which is sufficiently regular for measurement usually consists of two hexagonal pyramids joined base to base, one of the pyramids being truncated, and the angle formed by the junction of the pyramids being about 80 degrees; and F. Leydolt 33 has
- * Accounts of severe hailstorms, in some of which hailstones were reported as large as 13 to 20 inches in circumference, some weighing from one to four pounds, may be found in *Nature*, 125, 32, 656, 728, 765, 800, 840, 877, 913, 956, 994 (1930); 126, 41, 81, 117, 153, 188, 224, 262, 385, 457, 663, 669, 976, 1012 (1930); 137, 219-220 (1936).
 - 114 Dufour, L., Arch. des sci. phys. et nat. (N. S.), 10, 346-371 (1861).
 - 118 Berz, K. C., Kolloid Z., 41, 196-220 (1927).
 - 116 Blakiston, Captain, Proc. Rov. Soc. London, 10, 468 (1860).
 - 117 Secchi, P. A., Bull. Metcorol. Osserv. Coll. Romano, 15, 73-74 (1876).

 - 118 Humphreys, W. J., Monthly Weather Rev., 56, 314 (1928).
 119 Giblett, M. A., J. Roy. Acronaut. Soc. Grt. Brit., 31, 509-540-549 (1927).
 - 120 Grimminger, G., Monthly Weather Rev., 61, 198-200 (1933).
 - 121 Smithson, J., Ann. Phil. (N. S.), 5, 340 (1823).

reported that the optic axes of the constituent crystals in a hailstone are radial.

Descriptions of hailstones and of the microscopic appearance of sections of them have been published by J. H. L. Flögel. 121a

Snow and Frost.

Snow crystals are formed by inverse sublimation, by the passage of the molecules directly from the gaseous to the solid state (Hess). 65 , $^{9.}$ G. Tammann 121b states that snow is not formed at temperatures above -4 °C, that precipitation above -4 °C takes the form of rain. Obviously, the rain may freeze, producing hail. G. Stüve 121c has concluded that gaseous nuclei of condensation give rise to drops of water only; that soluble salts give rise to drops if the temperature at which the condensation begins is above -20 °C, and to stars of snow if the temperature is lower; and that insoluble hygroscopic nuclei give rise to needles of ice at all temperatures below 0 °C.

The typical crystal of snow or frost is 6-rayed, but innumerable modifications are found. The finest collection of photographs of snow-crystals and frost figures is that of W. A. Bentley, containing over 4000 negatives. He states that in the 45 years of his study he has never seen two snow crystals that were exactly alike. He has published an extended study of such crystals, ¹²² and recently, in conjunction with W. J. Humphreys, has published a beautifully illustrated volume (230 pages) entitled "Snow Crystals" (1931). Microphotographs of snow and rime have been published by G. Stüve ^{121c}; detailed studies of crystals of snow and frost, both natural and artificial, have been carried out by U. Nakaya and associates, ¹²³ and by G. Hellman, ¹²⁴ I. B. Schukewitsch, ¹²⁵ and A. Erman. ¹²⁶ Snow and the structure and properties of snow fields and the changes they undergo have been studied by G. Seligman, ^{73, 127} and G. Seligman and C. K. M. Douglas. ¹²⁸

When a thin layer of mud freezes during a cold night, the ice often takes the form of loosely packed hexagonal columns, often hollow, each carrying on its top a grain of sand or a bit of earth; or the entire bundle of columns may be covered with a continuous roof of earth. It is commonly stated that the optic axis of the ice in these columns is always vertical, ^{108, p. 12} but F. Klocke ⁷⁵ found this to be seldom true. He found that each column was generally an aggregate of small, approximately parallel needles meeting at sharp angles, and having their optic axes variously

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1218 Flögel, J. H. L., Ann. d. Physik (Pogg.), 146, 482-486 (1872).
121b Tammann, G., "Aggregatzustände," p. 219, 1922.
121c Stüve, G., Gerlands Beitr. zu Geophys. (Köppen Bd. 1), 32, 326-335 (1931).
122 Bentley, W. A., Monthly Weather Rev., 29, 212-214 (1901); 35, 348+, 397+, 439+, 512+, 584+ (1907).
123 Nakaya, U., and associates, J. Fac. Sci. Hokkaido, 1934-1936; see Seligman, G., Nature, 140, 345-348 (1937).
124 Hellman, G., "Schneekristalle," 1893.
125 Schukewitsch, I. B., Bull. Acad. Imp. Sci. St. Petersburg (6), 4, 291-302 (1910).
126 Erman, A., Phil. Mag. (4), 17, 405-413 (1859).
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Seligman, G., Nature, 140, 345-348 (1937); J. Roy. Meteorol. Soc., 63, 93-103 (1937).
 Seligman, G., and Douglas, C. K. M., "Snow Structure and Ski Fields," xii + 555, 1936.

directed, so that when placed in parallel light between crossed nicols it was impossible to orient a column in such a way as to obtain darkness. The composite character of the columns has been mentioned by Yoshida and Tsuboi also (see p. 409).

Glaciers.

Glaciers and the icebergs to which they give rise are merely compacted masses of snow and frost. They contain large amounts of entrapped air. H. T. Barnes 129 has found that the volume of the entrapped air, when under a pressure of 1 atmosphere and at the temperature of the iceberg, may be from 7 to 15 per cent of the total volume of the berg.

Icebergs derived from glaciers may explode with considerable violence, as a result of their internal strains. A. E. Nordenskjöld 130 likens them to immense Prince Rupert drops. He gives (pp. 319, 320) the following description of the breaking up of such bergs: "Glacier-ice shows a great disposition to fall asunder into small pieces without any perceptible cause. It is full of cavities, containing compressed air which, when the ice melts, bursts its attenuated envelope with a crackling sound like that of the electric spark. Barents relates that on the 20/10th August 1596 he anchored his vessel to a block of ice which was aground on the coast of Novaya Zemlya. Suddenly, and without any perceptible cause, the rock of ice burst asunder into hundreds of smaller pieces with a tremendous noise, and to the great terror of all the men on board. Similar occurrences on a smaller scale I have myself witnessed."

As the surface of the glacier melts, it becomes fissured and pitted, and acquires a granular structure. These "glacier grains" vary in size from that of snow crystals to several hundred cubic centimeters 108, pp. 166, 167: E. Hagenbach-Bischoff 62 says they may be 10 or even 15 cm broad. Each grain is a single crystal, although its boundaries are very irregular, and generally curved, and its refraction may be complicated by the strains that exist in it: Tyndall's flowers of ice are circular. 131 "Glacier ice is a sort of conglomerate of these grains, differing, however, from a conglomerate proper in that there is no matrix, the grains fitting each other perfectly." 132 It would, perhaps, be better to say that the matrix consists solely of imperceptibly thin layers separating the grains and enclosing each in a separate cell. (See p. 401 + .) S. Skinner has studied the fine structure of the surface-ice of glaciers by the use of plaster casts. 133

In general, the optic axes of the several grains seem to be arranged quite at random, 81a, 62, 181 but Drygalski has reported that the optic axes of the several grains in the old inland Antarctic ice-sheet and of those in the deeper layers of Antarctic icebergs are similarly directed (see G. Tammann 121b, pp. 214, 215). C. Grad, 184 and A. Bertin 82 have reported the same

¹²⁰ Barnes, H. T., "Ice Engineering," p. 346, 1928.

 ¹²⁰ Nordenskjöld, A. E., "The Voyage of the Vega around Asia and Europe," 1882.
 121 Klocke, F., Neues Jahrb. Mineral., Geol., 1881, 23-30 (1881).
 122 McConnell, J. C., and Kidd, D. A., Proc. Roy. Soc. (London), 44, 331-367 (1888).

¹⁸⁸ Skinner, S., Proc. Camb. Phil. Soc., 11, 33-36 (1901).

thing for the deeper portions of the termini of glaciers, the direction of the optic axis being vertical. Grad attributed this orientation to the prolonged action of pressure, and likened it to the temporary birefringence produced in glass by suitable stresses. Bertin seems to have thought that it is to be accounted for in somewhat the same way as the vertical direction of the optic axes of the crystals in a sheet of ice formed on calm water. O. Mügge ⁸¹ would explain it as a special case of his experiment in which a bar of ice supported near its ends and loaded in the middle, the optic axis being perpendicular to the length of the bar and inclined to the vertical, tends to rotate so as to bring the optic axis toward the vertical.

On the other hand, J. Müller ¹³⁵ found no such general uniformity in the directions of the axes, though in small portions of the ice, scattered here and there through the lower end of the glacier, the axes were vertical.

The grains increase in size, and consequently decrease in number, as the ice ages. This occurrence in the body of a glacier does not result from the freezing of water seeping down from the melting surface, for it has been found that the body of the glacier is impervious to liquids, and that at depths exceeding a meter the temperature is rarely as high as 0 °C (Vallot ⁹⁰ and Forel as quoted by Hagenbach-Bischoff ¹³⁶). It is probably due to migratory recrystallization (p. 412)—to a sublimation and recondensation similar to that by means of which large rain-drops devour smaller ones,^{74, 62} assisted by the tendency of crystals in close contact and under pressure to freeze together solidly if their axes are similarly directed (p. 413 and ^{62, 136}), and by the melting and refreezing that accompany stresses and their variations (p. 437 + and E. Hagenbach-Bischoff ¹³⁷).

But through the upper layers of the glacier, especially through the névé, water can percolate. Hence if the surface temperature during the day is above 0° C there will be melting, percolation of water, and freezing of that water at greater depths. Thus the observed internal layers of ice may be formed, the grains may grow, and possibly an actual glacier might be formed in this manner, ¹³⁸ but only in a relatively thin superficial layer can the growth of the grains be so affected. ¹³⁹

Those interested in the structure of a snow field and in changes it undergoes as it is gradually converted into a glacier, should read the papers by G. Seligman, 78, 127 and G. Seligman and C. K. M. Douglas. 128

A glacier flowing down a mountain is squeezed together where its bed narrows, exhibiting definite lines of flow. Where the bed widens the ice splits, forming crevasses; these form suddenly, but widen slowly.¹⁴⁰

¹³⁴ Grad, C., Compt. rend., 64, 44-47 (1867); see Heim, A., Ann. d. Physik (Pogg.) Erg. Bd., 5, 30-63 (1871).

¹⁸⁵ Müller, J., Ann. d. Physik (Pogg.), 147, 624-626 (1872).

¹⁸⁶ Hagenbach-Bischoff, E., Z. Kryst., 20, 309-310 (1892) ← Verh. d. naturf. Ges. Basel, 8, 821-832 (1889).

¹⁸⁷ Hagenbach-Bischoff, E., Z. Kryst., 11, 110-111 (1885) ← Verh. naturf. Ges. Basel, 7, 192-216 (1882).

¹²⁸¹ l)evaux, J., Compt. rend., 185, 1602-1604 (1927).
1280 See also, Emden, R., Noue Denkschr. allgem. Schweis Ges. ges. Naturwiss., 33 (1892).
140 Barnes, H. T., "Ice Formation," p. 92, 93, 1906.

The mechanics of glaciers has been discussed recently by M. Lagally.¹⁴¹ Proposed explanations of the motion of glaciers have given rise to various disputes, such as the recent one between R. T. Chamberlin ¹⁴² and O. D. Engeln.¹⁴³

Of the many phenomena that may be involved in the motion of glaciers, the one most commonly invoked to explain the general flow and the compression of the glacier where the bed narrows is that of the lowering of the melting-point by pressure, together with the regelation that follows a release of pressure. ¹⁴⁴ G. Beilby, ⁸⁸, ^{pp}. ¹⁹⁴⁻²⁰⁰ however, has pointed out that this can be effective only when the temperature of the ice is near 0 °C, a pressure of 138 atm (corresponding to a free column of ice 5000 feet high) causing a depression of only 1 °C in the melting-point.

In contrast to the great pressure required to lower the melting-point by a small amount, he found that ice at -11 °C can be flowed by a steel burnisher exerting a pressure not exceeding 30 or 40 lbs. per sq. in. (say, 2 atm). In order to lower the melting-point by 11 °C, the pressure would have to be of the order of 1500 atm, corresponding to a free column of ice 55,000 ft. (over 10 miles) high.

He concluded that "true molecular flow, which," under differential stresses, "occurs alike at the external and internal surfaces of crystalline aggregates, has, therefore, a wider and more fundamental relation to the phenomena of ice flow than fusion and regelation." J. H. Poynting's (1881) conclusions (p. 431) regarding the melting of ice by pressure to which the resulting water is not subjected, and R. W. Wood's (1891) experiments (p. 439) on the compression of ice in which lead pellets were embedded, are of interest in this connection.

J. Vallot ⁹⁰ has studied the variation in the temperature with the depth below the surface of a glacier. He found that the diurnal variation did not extend below one meter, nor the annual variation below 6 or 7 meters. At a depth of a meter the temperature rose above 0 °C only under exceptional conditions. The work was done on Mt. Blanc, at an elevation of about 4.3 km.

An explanation of the veined structure often observed in glaciers, based on the melting and refreezing of ice under the action of changing stresses, has been offered by W. Thomson (Lord Kelvin).¹⁴⁵

H. Hess ^{108, p. 14} has stated that lightning had never been known to strike a glacier. But is it not probable that he has recorded merely a lacuna in our observations?

For a recent summary of our knowledge of glaciers, see H. Hess. 146

Lagally, M., Gerlands Beitr. Geophys., Suppl. Bd., 2, 1-94 (Bibliog. 112) (1933).
 Chamberlin, R. T., Science (N.S.), 80, 526-527 (1934).

<sup>Engeln, O. D., Idem, 80, 401-403 (1934); 81, 459-461 (1935).
Thomson, J., Proc. Roy. Soc. (London), 8, 455-458 (1857).</sup>

¹⁴⁵ Thomson, W. (Lord Kelvin), Proc. Rov. Soc. London, 9, 209-213 (1858).

¹⁴⁶ Hess, H., Müller-Pouillets Handbuch der Physik, 11 ed., Vol. 5, pp. 355-397, 1928.

Sea-ice.

The ice formed from sea-water contains some salt. This salt differs in composition from that contained in the sea. For example, the ratio of the sulfates to the chlorides is greater in the ice than in the sea. In the process of freezing a selection is made, certain constituents of the salt are in a measure retained, while others are more completely eliminated. It is to be expected that the retained salt is not contained in the ice-crystals themselves, but lies in the boundaries between the crystals. In newly formed sea-ice the salt is quite uniformly distributed in the proportion of 4 or 5 parts per 1000 of ice. As the ice ages, there is a migration of the salt from the interior to the surface. In a case reported by Drygalski, the salinity decreased in two months from 4 or 5 parts per 1000 to only 1 or 2 (see H. T. Barnes 148), and A. E. Nordenskjöld 130 stated: "The water which is obtained by melting sea-ice is not completely free from salt, but the older it is the less salt does it contain" (p. 321).

J. Y. Buchanan ⁵³ has shown that the salt contained in sea-ice is not contained in the ice crystals, nor as a solid inclusion, but in brine entrapped between the crystals. When the ice is first formed, the composition of that salt is the same as that in the water from which the ice was formed, but as the ice ages, the composition changes, owing to various secondary effects. Owing to its composite structure—ice-crystals surrounded by brine—sea-ice melts progressively, the amount melted at any given temperature being just enough to make the concentration of the resulting intercrystallic brine such that the melting point of ice bathed in the brine is the given temperature.

In this composite structure and in the progressive melting is to be found the explanation of the many differences between the behavior of sea-ice and of ice formed from fresh water. For example, sea-ice not only melts below 0 °C, but as the temperature is raised from a low value, the volume of the ice reaches a maximum and then decreases as the temperature is further increased. This decrease indicates that melting has already begun, although the ice appears to be as solid as ever; the melting is at the boundaries of the crystals, where they are bathed with the intercrystallic brine. Thus, ice containing 2.73 parts of chlorine per 1000 began to contract at -14 °C, and that containing 6.49 parts began at -18 °C. Buchanan has shown that such contraction can be explained in the manner indicated. Pettersson 147 gives several sets of data showing such contraction.

Buchanan stated: "At the winter quarters of the Vega brine was observed oozing out of sea-water ice and liquid at -30 °C. It was very rich in calcium and especially magnesium chlorides. In fact, it is probably quite impossible by any cold occurring in nature to solidify sea-water."

Thin layers of sea-ice are white, from mechanically suspended salt, and

 ¹⁴⁷ Pettersson, O., "Vega-Expeditiones Vetensk. Jakt.," Vol. 2, pp. 249-323, 1883; as reviewed in Beibl. Ann. d. Physik, 7, 834-841 (1883).
 ¹⁴⁸ Barnes, H. T., "Ice Engineering," p. 231, 1928.

are so mobile that a small wave may travel through them without breaking them up.148

The following description of the formation of sea-ice is adapted from one that Barnes 148, p. 232 credits to J. B. Woodyatt. At first there appears a sort of thin slush on the surface of the sea. From a distance, its appearance resembles that of oil on water. It forms a cohering and flexible surface; the wash from a ship distorts the surface, but does not break it up. This slush forms into little discs about 4 inches in diameter, which gradually grow in diameter. They have no power of cohesion. They are pushed about by wind and water until several are piled partly on top of one another, making aggregates about 2 feet in diameter, the intervening spaces being filled with the slush. The lapping of the water deposits slush in ridges both on their tops and along their edges. Clumpets with such slushy edges tend to stick together when they meet, but even a small wave will pull them apart or slide one on top of another. Presently the intervening slush hardens, cementing the surface, then the ice grows rapidly.

At other times the growth proceeds quite differently, giving rise to much clearer and more brittle ice, with vertical cleavage planes. This type of ice forms on still water in very cold weather.

60. MOLECULAR DATA FOR ICE

Numerous suggestions regarding the nature and structure of the ice molecule have been advanced, but no generally accepted conclusion has yet been arrived at. Here the compiler will do no more than indicate a few of those suggestions, and cite certain publications in which the subject is discussed in some detail. The corresponding sections for the vapor (9) and the liquid (25) should be consulted.

Association of Molecules in Ice.

H. M. Chadwell 149 has reviewed the several suggestions regarding the molecular structure of water and of ice, and the evidence on which they rest. He gives a bibliography of over 100 titles.

The most widely held opinion seems to be that the molecule of ice-I is (H₂O)₃, called trihydrol, but several have regarded it as more complex, and G. B. B. M. Sutherland 150 concluded that nothing more complex than (H₂O)₂ is needed. To Sutherland's conclusions, I. R. Rao ¹⁵¹ seriously objects, favoring the trihydrol theory.

In contrast with the preceding, others, including W. H. Bragg 152 and W. H. Barnes, 158 are of the opinion that a mere space lattice of ions is preferable to any type of polymerization as a representation of the structure

¹⁴⁹ Chadwell, H. M., Chem'l Rev., 4, 375-398 (1927).

¹⁸⁰ Sutherland, G. B. B. M., Proc. Roy. Soc. (London) (A), 141, 535-549 (1933).

 ¹⁵¹ Rao, I. R., Idem, 145, 489-508 (1934).
 152 Bragg, W. H., Proc. Phys. Soc. (London), 34, 98-103 (1922).

¹⁸⁸ Barnes, W. H., Proc. Roy. Soc. (London) (A), 125, 670-693 (1929).

of ice-I. To that, T. M. Lowry and M. A. Vernon ¹⁵⁴ do not agree, giving reasons for believing that there is a polymerization in which additional bonds come into play. They postulate a network of single bonds between quadrivalent oxygen and bivalent hydrogen.

From his determinations of the dielectric constant of ice-I and of the way it varies with the temperature and the frequency, J. Errera ¹⁵⁵ thought it probable that at the freezing point there is no distinction between the molecules of water and of ice. That does not accord with the more common opinion that the proportion of trihydrol in water at 0 °C is less than 50 per cent of the whole (Table 79).

I. R. Rao ¹⁵¹ has expressed the opinion that it is not possible to derive from existing x-ray data any definite conclusion regarding the extent of the association in either ice or water.

The complexity of the ice molecule has been considered also by P. N. Chirvinskii,¹⁵⁶ J. Duclaux,¹⁵⁷ R. de Forcrand,¹⁵⁸ E. J. M. Honigmann,¹⁵⁹ and L. Schames,¹⁶⁰

From the great values of the pressure-derivative of their mutual equilibrium temperatures (small values of dP/dt, Table 270), G. Tammann ¹⁶¹ has concluded that ices I, III, V, and VI have all the same molecular weight, their molecules being isomeric, differing one from another in the distances between the constituent atoms, but not in the grouping of them.

Structure of the Molecule of Ice.

The structure of the molecule of ice, as regards the arrangement of the atoms, the distances between them, and their bonding, is intimately related to the ultimate crystalline structure of ice, as revealed by means of x-rays. Both will be considered in this section, the second being considered first. For the actual values of the periodicities observed, see Table 212. H. M. Chadwell's review, 140 with bibliography, should be consulted.

Only ice-I, ice-II, and ice-III have as yet been studied by x-rays. R. L. McFarlan has concluded that the lattice pattern of ice-II is that of a side-centered orthorhombic cell having a=7.80A, b=4.50A, and c=5.56A, and containing 8 molecules ¹⁶²; and that that of ice-III is a body-centered orthorhombic cell having a=10.20A, b=5.87A, and c=7.17A, and containing 16 molecules.¹⁶³

The x-ray studies of ice-I are numerous and conflicting (see A. B. Dobrowolski 164). A key to the last may perhaps be found in the recent

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    Lowry, T. M., and Vernon, M. A., Trans. Faraday Soc., 25, 286-291 (1929).
    Errera, J., J. de Phys. (6). 5, 304-311 (1924).
    Chirvinskii, P. N., Chem. Abst., 17, 2525 (1923) ← Bull. Soc. Russe amis l'étude l'univers (Petrograd), 7, 6-10 (1918).
    Duclaux, J., J. de chim. phys., 10, 73-109 (1912).
    6 Forcrand, R., Compt. rend., 140, 764-767 (1905).
    Honigmann, E. J. M., Naturwissenschaften, 20, 635-638 (1932).
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¹⁰⁰ Schames, L., Ann. d. Physik (4), 38, 830-848 (1912).

¹⁰¹ Tammann, G., "Aggregatzustände," pp. 143, 144; 1922.
102 McFarlan, R. L., J. Chem'l Phys., 4, 60-64 (1936) → Phys. Rev. (2), 49, 199 (A) (1936).
103 McFarlan, R. L., J. Chem'l Phys. 4, 253-259 (1936) → Phys. Rev., 49, 644 (A) (1936).

announcement by N. Seljakov ¹⁶⁵ that ice-I occurs in two forms: α -ice, belonging to one of the following 4 classes, 3 of the hexagonal system—dihexagonal-bipyramidal (D_{6h}) , trapezohedral (D_6) , and dihexagonal-pyramidal (C_{6v}) —and one—the ditrigonal-bypyramidal (or holohedral) class (D_{8h}) —of the trigonal system; and β -ice, belonging either to the rhombohedral class (C_{8i}) or the pyramidal class (C_3) of the trigonal system. The lattice unit is essentially the same for each form: $a=4.52\pm0.03$ A, $c=7.34\pm0.04$ A, $c/a=1.60\pm0.02$. He discusses the discrepancies in the reported values in the light of his observations.

A recent x-ray analysis of ice-I has been made by W. H. Barnes, ¹⁵³ using both single crystals obtained from commercial, artificially frozen ice and thin plates of clear, flawless ice grown on the surface of a basin of water exposed to the air during cold weather. He obtained Laue photographs of the plates at -78.5 °C, and rotation and oscillation photographs of the single crystals at -20 °C. The former showed that the crystals can be referred to hexagonal axes, and the latter gave for the unit cell:

$$a = 4.53_5$$
A, $c = 7.41$ A, $c/a = 1.634$

and content = 4 hydrol (H₂O) molecules. The dimensions are believed to be correct within a few parts in 1000.

These results essentially agree with those (a = 4.52, c = 7.32, c/a = 1.62) of D. M. Dennison ¹⁶⁶ for ice obtained by plunging into liquid air a capillary tube filled with water. Dennison's data are given in R. W. G. Wyckoff's compilation ¹⁶⁷ together with the citations: W. H. Bragg, ¹⁵² R. Gross, ¹⁶⁸ F. Rinne, ¹⁶⁹ and A. St. John, ¹⁷⁰

On the other hand, A. St. John,¹⁷⁰ using single crystals obtained by freezing water in exposed open pans, found $a=4.74\Lambda$, from which he computes $c=6.65\Lambda$, accepting Dana's value c/a=1.4026. Both H. T. Barnes ^{148, p. 20} and W. H. Barnes ^{153, p. 672} suggest that the difference between these and the other values may arise from a real difference in the structure of the specimens used.

Laue photographs of powdered ice at -9, -13, -78, and -183 °C were taken by W. H. Barnes, ^{153, p. 672} but no indication that the structure depends at all upon the temperature was found.

On the other hand, E. F. Burton and W. F. Oliver 171 have found that the structure of the ice formed by freezing water-vapor onto a cold metal surface below -80 °C varies with the temperature, becoming vitreous if the temperature is below about -110 °C. If the temperature of the vitreous ice is raised above -110 °C its structure becomes more organized, but

¹⁶⁴ Dobrowolski, A. B., Bull. Soc. Fr. Mineral., 56, 335-346 (1933).

¹⁶⁵ Seljakov, N., Compt. rend. Acad. Sci. URSS, 10, 293-294 (1936); 11, 227 (1936); 14, 181-196 (1937).

¹⁰⁰ Dennison, D. M., Phys. Rev. (2), 17, 20-22 (1921).

¹⁶⁷ Wyckoff, R. W. G., Int. Crit. Tables, 1, 338-353 (341) (1926).

¹⁶⁸ Gross, R., Centralbl. Min., Gcol., Palaon, 1919, 201-207 (1919).

¹⁰⁰ Rinne, F., Ber. Sächs. Ges. Wiss. (Math.-Phys.), 69, 57-62 (1917).

¹⁷⁰ St. John, A., Proc. Nat. Acad. Sci., 4, 193-197 (1918).

¹⁷¹ Burton, E. F., and Oliver, W. F., Proc. Roy. Soc. (London) (A), 153, 166-172 (1935).

does not reach the normal structure before -80 °C is reached (see Table 212).

From his own observations, D. M. Dennison ¹⁶⁶ inferred a close-packed hexagonal lattice consisting of two sets of interpenetrating prisms; whereas W. H. Barnes ^{153, p. 672} concluded from his that the structure is either ditrigonal bipyramidal (D_{3h}^4) or dihexagonal bipyramidal (D_{6h}^4) , with the probabilities in favor of the latter, and that an ionic structure is to be preferred. He proposed the D_{6h}^4 structure with an H placed at the middle of each line joining a pair of adjacent O's. He essentially agrees with W. H. Bragg, ¹⁷² who regards the molecular structure of ice as hexagonal and differing from that of diamond simply by the replacement of each C by an O, and the insertion of an H between the members of each pair of O's. Such a structure imposes on O a covalence of 4 and on H that of 2. Some of the conclusions that flow from the assumption of such covalences have been discussed by S. W. Pennycuick. ¹⁷³ See also T. M. Lowry and M. A. Vernon, ¹⁵⁴ and R. de Forcrand. ¹⁵⁸

On the other hand, E. L. Kinsey and O. L. Sponsler ¹⁷⁴ infer a different structure from the same observations by W. H. Barnes, ^{153, p. 672} one in which occur the units H^+ and $(H_3O_2)^-$, the latter having the form of a double tetrahedron with the O's at the extremities of the axis.

W. H. Bragg ¹⁷² has concluded that the distance between the centers of adjacent O-atoms is 2.76A, between neighboring atoms lying in the same plane is 4.52A, and between consecutive basal planes is 3.67A; and M. L. Huggins, ¹⁷⁵ that the radius of the H-atom in ice, defined as "the distance from nucleus to valence electron-pair," is 0.73A.

Reasons have been presented for believing that each O-atom in ice is tetrahedrally surrounded by 4 others, and that each H-atom lies on the line connecting 2 adjacent O-atoms, but nearer to one of those atoms than to the other. I. Pauling ¹⁷⁶ gives these distances as 0.95A and 1.81A; and P. C. Cross, J. Burnham, and P. A. Leighton ¹⁷⁷ give them as 0.99A and 1.77A.

That the structure of ice is like that of tridymite was proposed by J. D. Bernal and R. H. Fowler ¹⁷⁸ (see p. 174), and seems to be widely accepted. But W. H. Barnes ¹⁷⁹ has cautioned against a too hasty or uncritical acceptance of that proposal, doubting if any modification of the structure proposed by himself was yet necessary. See also, M. L. Huggins, ¹⁸⁰ L. Pauling, ¹⁷⁶ W. F. Gaiuque and J. W. Stout. ¹⁸¹

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    Bragg, W. H., Proc. Phys. Soc. (London), 34, 98-103 (1922).
    Pennycuick, S. W., J. Phys'l Chem, 32, 1681-1696 (1928).
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¹⁷⁴ Kinsey, E. L., and Sponsler, O. L., Phys. Rev. (2), 40, 1035-1036 (A) (1932); Proc. Phys. Soc. (London), 45, 768-779 (1933).

¹⁷⁶ Huggins, M. L., Phys. Rev. (2), 21, 205-206 (1923).

¹⁷⁰ Pauling, L., J. Am. Chem. Soc., 57, 2680-2684 (1935) → Nature, 137, 327 (1936).

²⁷⁷ Cross, P. C., Burnham, J., and Leighton, P. A., J. Am. Chem. Soc., 59, 1134-1147 (1937).

Bernal, J. D., and Fowler, R. H., J. Chem'l Phys., 1, 515-548 (1933).
 Barnes, W. H., Trans. Roy. Soc. Canada III (3), 29, 53-59 (1935).

¹⁸⁰ Huggins, M. I., J. Am. Chem. Soc., 58, 694 (L) (1936); J. Phys'l Chem., 40, 723-731 (1936).

¹⁸¹ Giauque, W. F., and Stout, J. W., J. Am. Chem. Soc., 58, 1144-1150 (1936).

61. Interaction of Ice and Corpuscular Radiation

No data have been found regarding either the absorption of corpuscular radiation by ice (range of α -rays, absorption of β -rays, etc.) or the effect of such radiation upon ice (electron emission excited by +ions, by electrons, etc.); but T. H. Johnson ¹⁸² has shown that atomic beams (hydrogen) are reflected from ice in preferential directions.

62. Adhesiveness of Ice

J. W. McBain and D. G. Hopkins ¹⁸³ have reported that the freezing of a thin film of water between two plates of fused silica produces a joint that is "very strong" in shear. The freezing was done with solid CO₂, and the test was presumably made at that temperature.

63. SLIDING FRICTION OF ICE

J. Joly ¹⁸⁴ seems to have been the first to point out that the slipperiness of ice and the "biting" of a skate are to be explained by the melting of ice under pressure. As the ice is melted, the curved runner of the skate sinks into the ice until the bearing surface becomes of such a size that the load is just insufficient to lower the melting point below the temperature of the surroundings. The sinking of the runner gives rise to the "bite," and the layer of water, acting as a lubricant, causes the slipperiness. When the ice is very cold, the sinking will be slight, and, as is well known, the "bite" is poor; then, hollow-grinding is of advantage, as it reduces the area of the bearing surface, thus increasing the pressure. The same explanation of the slipperiness of ice was advanced later by O. Reynolds. He stated that Nansen, "in his book on Greenland," says that at very low temperatures ice completely loses its slipperiness.

When one solid body slides over another, the coefficient of friction (f) and the angle of repose ϕ are related as follows to the normal force (L) with which one body is pressed against the other and the tangential force (F) required to slide one body over the other with a constant velocity: F = fL, $\tan \phi = f$, where f is independent of L, unless that is so great as to distort the surfaces; and ϕ is the angle at which the surface of contact must be inclined to the horizontal if the upper body, when once started, slides with constant velocity, F and L arising solely from the action of gravity.

Applying the preceding to the friction between ice and aluminum at a constant temperature near -6.5 °C, H. Morphy ¹⁸⁶ has found two distinct values for f, depending upon the load; the bearing surfaces were small, but of unknown area. If the load was under 14.3 g*, then $f = 0.36 \pm 0.01$,

¹⁸² Johnson, T. H., Nature, 120, 191 (1927).

¹⁸⁸ McBain, J. W., and Hopkins, D. G., Dept. Sci. Ind. Res. (Gt. Brit.), "2nd Report Adhesives Res. Com.," pp. 34-89 (41) (1926).

¹⁸⁴ Joly, J., Proc. Roy. Dublin Soc. (N. S.), 5, 453-454 (1886).

 ¹⁸⁵ Reynolds, O., "Papers on Mechanical and Physical Subjects," Vol. 2, pp. 734-738, Cambridge Univ. Press, 1901; ← "Mem. and Proc. Manchester Lit. Phil. Soc.," Vol. 43, 1899.
 186 Morphy, H., Phil. Mag. (6), 25, 133-135 (1913).

 ϕ about 20°; if the load exceeded 15 g*, $f = 0.17 \pm 0.01$, ϕ about 9.5°. Within each range, f was independent of the load.

The inverse case, i.e., ice sliding down an inclined plane, has been studied by W. Hopkins ¹⁸⁷; and H. Moseley ^{187a} reported that he had repeated Hopkins's experiments, with many modifications, and had verified his conclusions.

Using a rough-hewn slab of sandstone so mounted that it could be rotated about an axis parallel to its plane and perpendicular to the grooves made by the tool, Hopkins found that the angle of repose for a block of polished marble resting on it was 20° . Ice loaded to a pressure of 150 lb*/ft² slid down the slab at the following unaccelerated rates:

Inclination (°)	3	6	9	12	15
Rate (in/hr)	0.31	0.52	0.96	2.0	2.5

At an inclination of 20° the motion was accelerated. At an inclination of 9° the removal of 2/3 of the load reduced the rate by nearly half. Even at 1° there was a perceptible motion. On a smooth, but unpolished, slab of the same kind of stone there was a perceptible motion when the inclination was only 40′, and on polished marble there was motion at the "smallest possible inclination. The motion, in fact, afforded almost as sensitive a test of deviation from horizontality as the spirit-level itself."

In all these cases the ice "melted continuously but very slowly" at its surface in contact with the slab. He attributed the unaccelerated motion "to the circumstance of the lower surface of the ice being in a state of constant, though slow distintegration." The water acted as a lubricant.

When there was no melting, the angle of repose on sandstone was about the same for ice as for marble, and about equal to the inclination (20°) for the initiation of accelerated motion when there was melting.

64. DEFORMABILITY OF ICE

The ways in which the volume and form of a block of ice vary with its uniform temperature and with the uniform hydrostatic pressure to which it is subjected will be found in Section 67. Such changes in form are slight, depending solely upon the anisotropy of the block.

Changes produced by non-uniform stresses, and the way they vary with the temperature, will be considered here. Under such stresses, the form of a block may be changed enormously.

Although a large number of experiments of various kinds bearing upon the behavior of ice under non-uniform stresses have been reported, only a few of them are of such a kind as to yield numerical data of general applicability. Others, however, are of much interest, and contribute valuable descriptive information regarding the characteristics of ice. For these reasons they deserve a place in such a compilation as this. The experiments cannot be satisfactorily separated into mutually exclusive groups.

¹⁸⁷ Hopkins, W., Phil. Mag. (3), 26, 1-16 (1845). ^{187a} Moseley, H., Idem (4), 42, 138-149 (1871).

They will be considered under the following heads, those yielding no numerical data of general applicability being termed descriptive, even though many of them contain quantitative data.

Descriptive treatment Quantitative treatment Young's modulus Linear compression Extension Poisson's ratio Flexure Rigidity Tensile strength Punching Penetration Strength, linear compression Flowing Shearing strength Recovery Hardness Brittleness Plasticity and viscosity Sustaining power

Whatever other phenomena may be involved, most permanent deformations produced in ice by non-uniform stresses involve a melting where the stress is great, the flowing of the supercooled water thus produced, and the refreezing of that water when its pressure has been reduced—all in accordance with the conclusions reached by J. Thomson.¹⁸⁸

It should be remembered that such melting can occur only when the melting point of ice under the stress is lower than the existing temperature of the ice, and also, as shown in the second of Thomson's articles, that when the stress is borne by the ice alone such positive melting will occur whatever the nature of the stress—pressure, tension, torsion, etc.—and the attendant lowering of the temperature will lead to the formation elsewhere of unstressed ice from the melt, except as such formation is impeded by the absence of nuclei suitable for its initiation. See also E. Riecke, 189

On account of such effects, a mass of ice might flow and exhibit plasticity although its individual crystals remained unbroken and perfectly elastic.

The early papers treating of melting under pressure and of regelation should be read, *i.e.*, those already cited and J. Thomson, ¹⁹⁰ W. Thomson (Lord Kelvin), ¹⁹¹ M. Faraday, ¹⁹² and the topic Regelation in this volume (p. 412) should be consulted.

J. Johnston and L. H. Adams ¹⁰³ have given reasons for believing "that every *permanent* deformation of a crystalline aggregate is conditioned by, and consequent upon, a real melting," which melting is in general to be ascribed to an inequality in the pressures on the liquid and on the solid.

¹⁸⁸ Thomson, J., Trans. Roy. Soc. Edinburgh, 16, 575-580 (1849); Proc. Roy. Soc. (London), 11, 473-481 (1861).

¹⁸⁹ Riecke, E., Ann. d. Physik (Wied.), 54, 731-738 (1895).

¹⁹⁰ Thomson, J., Proc. Roy. Soc. (London), 8, 455-457 (1857); 10, 152-160 (1859); 11, 198-204 (1861).

¹⁹¹ Thomson, W. (Lord Kelvin), Phil. Mag. (3), 37, 123-127 (1850); Proc. Roy. Soc. (London), 9, 141-143, 209-213 (1858).

¹⁰² Faraday, M., Proc. Roy. Soc. (London), 10, 440-450 (1860); "Exp. Res. in Chem. and Phys.," pp. 372-374, 377-382, 1859.

¹⁹⁸ Johnston, J., and Adams, L. H., Am. J. Sci. (4), 35, 205-253 (211) (1913).

They accept J. H. Poynting's conclusion ¹⁹⁴ that, when ice is subjected to a pressure of P atm more than that to which the water in contact with it is subjected, the melting point is lowered by 0.0862P °C, 11.5 times as many degrees as it would have been if this additional pressure had been exerted upon the water as well as the ice. In that paper, Poynting gives experimental evidence that such is the case. Johnston and Adams state that such a lowering is accepted by Roozeboom, Ostwald, LeChatelier, and Nernst, but not by G. Tammann. ¹⁹⁵ Whether, and in how far, such melting accounts for the ease with which ice at -11 °C was flowed by G. Beilby ¹⁹⁶ remains to be determined (see p. 422).

Both J. Y. Buchanan ¹⁰⁷ and G. Quincke ¹⁹⁸ were of the opinion that the mechanical deformation of ice is greatly facilitated, if not conditioned, by the presence between the crystals of thin films of liquid (a solution) of low melting point. They believed that such films are always present.

Discussions of the deformability of ice may be found in such treatises as those of G. Tammann, ¹⁹⁹ H. Hess, ¹⁰⁸ G. Beilby, ⁸⁸ and H. T. Barnes, ²⁰⁰ as well as in the scientific journals.

Descriptive Treatment.

Lincar Compression.—In his abstract of the paper which he presented before the Royal Society of London, Dec. 17, 1857, J. Tyndall ⁶⁰ described thus the behavior of a cylinder of ice subjected to longitudinal pressure. The cylinder "was placed between two slabs of boxwood and subjected to a gradually increasing pressure. Looked at perpendicular to the axis, cloudy lines were observed drawing themselves across the cylinder. Looked at obliquely, these lines were found to be the sections of dim surfaces which traversed the cylinder, and gave it the appearance of a crystal of gypsum whose planes of cleavage had been forced out of optical contact by some external force.

"The surfaces were not of plates of air, for they are formed when the compressed ice is kept under water. They also commence sometimes in the center of the mass, and spread gradually on all sides till they finally embrace the entire transverse section of the cylinder. A concave mirror was so disposed that the diffuse light of day was thrown upon the cylinder while under pressure. The hazy surfaces produced by the compression of the mass were observed to be in a state of intense commotion, which followed closely upon the edge of the surface as it advanced through the solid. It is finally shown that these surfaces are due to the liquefaction of the ice in planes perpendicular to the pressure.

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    Poynting, J. H., Phil. Mag. (5), 12, 32-48 (1881).
    Tammann, G., Ann. d. Physik (4), 7, 198-224 (1902); "Krystallisieren und Schmelzen," pp. 173-181, 1903.
    Beilby, G., "Aggregation and Flow of Solids," pp. 194-200, 1921.
    Buchanan, J. Y., Nature, 35, 608-611 (1887); 36, 9-12 (1887).
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¹⁹⁸ Quincke, G., Proc. Roy. Soc. (London) (A), 76, 431-439 (1905) → Nature, 72, 543-545 (1905).

<sup>Tammann, G., "Aggregatzustände," 1922.
Barnes, H. T., "Ice Formation," 1906; "Ice Engineering," 1928.</sup>

"The surfaces were always formed with great facility parallel to those planes in which the liquid flowers [flowers of ice] already described are produced by radiant heat, while it is exceedingly difficult to obtain them perpendicular to those planes. Thus, whether we apply heat or pressure, the experiments show that ice melts with peculiar facility in certain directions." (Cf. W. Thomson (Lord Kelvin).²⁰¹

Whence we may conclude that his description of the appearance and the formation of the surfaces applies primarily to cylinders so cut that the length is parallel to the optic axis of the ice; and that, under the action of linear compression along the axis, fusion occurs in discrete planes which are perpendicular to the axis.²⁰²

The intense commotion observed by Tyndall is probably related quite closely to that reported by Quincke 198 as occurring at the edge of an enlarging flower of ice (q.v.), and which he attributes to "a periodic capillary spreading out (Ausbreitung) of the salt solution of the foam-walls at the boundary between pure water and air or vacuum," except that in Tyndall's experiment a diffusion, rather than a capillary, phenomenon is probably involved.

In 1885, Koch reported that when subjected to a constant pressure (19 kg*/cm²) along its axis a certain cylinder of ice shortened at the following rates: At -5.7 °C, 0.9×10^{-4} per hr; at -2.5 °C, 17×10^{-4} per hr; at -0.9 °C, 126×10^{-4} per hr. The direction of the optic axis is not stated, but it was probably parallel to the axis of the cylinder.²⁰³

Three years later, McConnel and Kidd reported that blocks cut from a uniform sheet of ice and subjected to a pressure of 3.7 kg*/cm² perpendicular to the optic axis appeared to yield at the rate of only 0.1×10^{-4} per hr, but that even this was probably entirely spurious.²⁰⁴ The temperature seems to have been near 0 °C. The observations extended over 4 days. Even if this apparent yielding were true, and if the yielding were proportional to the pressure, these blocks under 19 kg*/cm² would yield only 0.5×10^{-4} per hr, as compared with the 126×10^{-4} per hr observed by Koch when the pressure was probably parallel to the axis.

For 3 pieces of glacier ice under a pressure of 3.2 kg*/cm^2 the yielding was 3.5×10^{-4} , 5.6×10^{-4} , and 0.7×10^{-4} per hr, respectively; the optic axes of the grains were randomly oriented. Observations extended over 5 days.

Similar observations have been reported by von Engeln.²⁰⁵

Extension.—The presence of numerous crevasses in every glacier, and the suddenness with which they are frequently formed, led students of glaciers to conclude that ice can yield to tension only elastically or by frac-

²⁰¹ Thomson, W. (Lord Kelvin), Proc. Roy. Soc. (London), 9, 141-143, 209-213 (1858).
2028 See also Tyndall's "The Forms of Water in Clouds and Rivers, Ice and Glaciers," New York, D. Appleton & Co., 1872.

²⁰⁸ Koch, K. R., Ann. d. Physik (Wied.), 25, 438-450 (1885).

²⁰⁴ McConnel, J. C., and Kidd, D. A., Proc. Roy. Soc. (London), 44, 331-367 (1888).

²⁰⁵ von Engeln, O. D., Am. J. Sci. (4), 40, 449-473 (1915).

ture; and this, in turn, gave rise to many experimental investigations of the behavior of ice under tension.

Observations of Pfaff 208 and the more extended ones of Fabian 207 showed that ice yields progressively when subjected to a continuing, constant tension. Main 208 stated: "Ice subjected to tension stretches continuously by amounts which depend on the temperature and the tensile stress. When the stress is great and the temperature not very low, there are extensions amounting to 1 per cent of the length per day. So continuous and definite is the extension that it can even be measured from hour to hour. These extensions took place at temperatures which preclude the possibility of melting and regelation. The extension increases continuously with all stresses above 1 kilo per square cm, and at all temperatures between -6 °C and freezing." He, and also Fabian, used bars of ice which had been frozen in a mold, and which, therefore, were probably conglomerates of crystals very variously oriented.

McConnel and Kidd 204 observed that not only the rate, but even the existence, of a progressive elongation depends upon the structure of the ice. They carried out extensive experiments on the stretching of bars cut in specified orientations from uniform sheets of ice. (By a uniform sheet is meant one in which the optic axis has the same direction at every point. They speak of using single crystals, but their report contains nothing to indicate that their crystallographic tests sufficed to distinguish between a single crystal and a homogeneous sheet containing many crystals similarly oriented as regards their optic axes. Cf. Mügge.) The work was extended by experiments on the bending of bars, by McConnel 45 and by Mügge 81 (p. 434).

They found that when the optic axis is perpendicular to the line of tension there is no measurable progressive stretching even when the stress has half the breaking value and the temperature is near 0 °C. But if the optic axis is inclined 45° to the line of tension there is a marked progressive stretching. There seem to have been no measurements on bars in which the optic axis is parallel to the line of tension. Bars of glacier ice, grains varying in diameter from 2 mm to 30 or 100 mm, stretched rapidly, as did mechanically molded ice of which the structure was surely very irregular. "The change in the rate of extension produced by an alteration in the tension, was in every case altogether out of proportion to the magnitude of the latter."

They found that an icicle, which "is an example of ice formed of very minute crystals irregularly arranged," (cf. p. 417, Icicles) stretched very slowly indeed. They were loath to ascribe the slow stretching to the multicrystalline structure, but later experiments by McConnel 209 convinced him

²⁰⁰ Pfaff, F., Ann. d. Physik (Pogg.), 155, 169-174 (1875); Sitz-ber. phys.-med. Soc. Erlangen, 7, 72-77 (1875).

 ²⁰⁷ Fabian, O., Rep. Exp.-Physik (Carl's), 13, 447-457 (1877) from Sitz.-Ber. Krakauer Akad.
 Wiss. (Math.-Nat. Kl.), 4, → Hess, H. 108, pp. 23, 28
 208 Main, J. F., Proc. Roy. Soc. London, 42, 329-330, 491-501 (1887).

²⁰⁰ McConnel, J. C., Prov. Roy. Soc. London, 49, 323-343, Exp. 8 (1891).

that the presence of intercrystalline faces does hinder plastic flow "by fettering the sliding of the layers in the separate crystals."

Flexure.—Reusch 200a reported, in 1864, that he had given a thin strip of ice a permanent set by bending it carefully by hand; and that a bar of ice, 10 by 1.2 cm by 0.3 cm thick, suspended horizontally by two slings 8 cm apart, and loaded in the middle with 180 grams, became visibly bent in 20 to 30 minutes, and the depression of its middle section increased to 6 or 8 mm before the bar broke. The temperature of the room was a few degrees above 0 °C, and the length and the breadth of the bar were probably parallel to the plane of freezing; that is, the optic axis was probably vertical.

Since then, many experiments have been made upon the progressive bending of bars of ice supported horizontally on narrow blocks near their ends, and loaded midway between the blocks.

From such experiments, Pfaff ²⁰⁶ concluded that near its melting point ice behaves like wax, the continued application of a force, no matter how small, producing a permanent deformation. Nothing is said regarding the direction of the optic axis, but it was probably vertical.

McConnel and Kidd ²⁰⁴ seem to have been the first to study the behavior of bars cut from ice of known uniform structure and in a specified direction with reference to the crystalline axes.

From their observations, and from other observations on bending, made by himself, McConnel ⁴⁵ concluded that "a crystal [of ice] behaves as if it were built up of an infinite number of indefinitely thin sheets of paper fastened together with some viscous substance which allows them to slide over each other with considerable difficulty; the sheets are perfectly inextensible and perfectly flexible. Initially they are plane and perpendicular to the optic axis; and when by the sliding motion they become bent, the optic axis at any point is still normal to the sheet at that point. Thus, when a bar with the optic axis transverse to its length is placed so that the axis is horizontal, and the sheets of paper consequently vertical and longitudinal, it refuses to take any plastic bend, however long the weight be applied. If the bar be now turned over, so that the sheets of paper are horizontal, quite a short interval suffices to produce a decided permanent depression of the middle of the bar."

Similar experiments by Mügge,⁸¹ Tammann,^{195, 1st} and Tammann and Salge ²¹⁰ confirmed and complemented McConnell's conclusions. As a result it may be stated that when the optic axis is horizontal and perpendicular to the length of the bar no permanent bend is produced, however long the load is applied. But if the same rod is rotated about its length so that the optic axis is vertical, a very short interval suffices to produce a decided permanent depression (McConnel, Mügge). If the bar is placed midway between these two positions, so that the optic axis, perpendicular to the length, is inclined 45° to the vertical, and the ends are so clamped that they cannot rotate about the longitudinal axis, then the application

²⁰⁰a Reusch, E., Ann. d. Physik (Pogg.), 121, 573-578 (1864).

²¹⁰ Tammann, G., and Salge, W., Neues Jahrb. Mineral., Geol., Beilage Bd., 57A, 117-130 (1928).

of the load causes the portion between the supports to bend and to rotate about the longitudinal axis in such direction as to place the optic axis more nearly vertical (Mügge). In all these cases the optic axis at any point of the deformed bar occupies the same position in the transverse section of the bar as it did before the bar was deformed.

"When the optic axis was longitudinal, the bar bent indeed, but not very readily, and the general behaviour was more obscure" (McConnel). This case involves a punching effect (see below).

By experiments on a rod cut from ice in which the planes of freezing were inclined to the optic axis by 50°, McConnel 45, 2nd (Experiment 14) showed that the phenomena just described are indeed related to the direction of the optic axis, and not to that of the planes of freezing. He concludes (Experiment 8) that the presence of interfaces between crystals hinders plastic flow "by fettering the sliding of the layers in the separate crystals."

Although somewhat differently pictured by him, Mügge's observations indicate that the sheets imagined by McConnel are permanently deformable without change in area, the permanence of the deformation being, perhaps, imposed by the "viscous substance" that binds the sheets together, rather than by the nature of the sheets themselves.

More recent, but apparently less extensive, work by M. Matsuyama ^{210a} confirms in the main those observations of McConnel and of Mügge upon which it touches, but Matsuyama concludes that the bending of such rods depends upon the relative displacements of the individual crystals rather than upon the distortions of the crystals themselves.

Such a picture of the structure of an ice crystal as was proposed by McConnel and extended by Mügge, supplemented by Tyndall's observation (1858)⁶⁰ that compression along the optic axis causes liquefaction in planes perpendicular to the axis, is of great value in any attempt to interpret observations having to do with the deformation of ice. (See also M. Faraday ⁵⁵ and J. Thompson.²¹¹)

Punching.—(See also Shearing strength, p. 449) O. Mügge 81 observed that when a bar of ice, cut with its length parallel to the optic axis, is supported horizontally on two blocks and most of the portion between the blocks is loaded uniformly by means of weights suspended from a stirrup resting on the top of the bar and nearly as broad as the distance between the blocks, then the portion directly under the stirrup is gradually forced downward without change in the direction of its optic axis. It is, in effect, punched from the bar. The entire periphery of the punched out portion is frequently marked by horizontal lines, but the ice remains perfectly clear, and exhibits no sign of cracks nor of optical anomalies due to strain. However the bar was rotated about the optic axis, no significant difference in the ease with which it can be punched was found; neither did a change in temperature from -3° to -16° C produce any marked effect.

snoa Matsuyama, M., J. Geol., 28, 607-631 (1920).

an Thomson, J., Proc. Roy. Soc. London, 11, 198-204 (1861).

There seems to be a minimum load below which punching does not occur. In one case a 5 kg* load produced no observable effect in 24 hrs, but when the load was increased to 7 kg*, the deformation was rapid.

When the length of the bar is perpendicular to the optic axis there is no such punching, however the axis may be oriented in the vertical plane.

When a portion is so punched from a bar whose length is parallel to the direction of the optic axis, the extent to which the ice has been punched from its initial position decreases progressively as one passes from either edge of the stirrup to the neighboring supporting block. Except possibly in extreme cases, the optic axis in these portions also remains horizontal. These lateral portions are marked by a series of faulting planes.

G. Tammann and W. Salge,²¹⁰ using a narrow stirrup to transmit the transverse punching force, have found that a longitudinal pressure of a few kilograms per cm² parallel to the optic axis may increase the number of these faulting planes some ten-fold if the temperature is -1 °C, but causes essentially no change in the number if the temperature is as low as -6 °C. If z is the punching force that must be applied in order to cause these planes to appear, to initiate a true punching, then at a fixed temperature. $\log_{10}(z_0/z) = bb$, b being the axially directed pressure; b is positive,* an increase in p producing a decrease in s. The value of s_0 depends upon instrumental details as well as upon the temperature. For certain conditions, they found the following values, the unit of z being 1 kg*, of p being 1 kg*/cm²: t = -1 °C, $z_0 = 1.5$, b = 0.52; t = -6 °C, $z_0 = 2.2$, b = 0.520.34; t = -12 °C, $z_0 = 2.8$, b = 0.11. These values indicate that both z_0 and b are linear in t; $z_0 = 1.4(1 - 0.086t)$, b = 0.56(1 + 0.066t). The lower the temperature the greater is the force required to initiate punching and the less is the effect of axially directed pressure.

Penetration.—If a solid object, such as a metal rod, tube, or ball, is pressed normally against a surface of ice, the pressure being maintained continuously, it gradually sinks into the ice, and the immediately surrounding ice rises in the form of a hillock. The rate at which a given object sinks under the action of a fixed pressure decreases as the temperature is reduced. Thus Pfaff 206 reports a case in which the object sank 3 mm in 2 hrs when the temperature was between -1 and 0 °C, 1.25 mm in 12 hrs when -4 to -3 °C, and, with 2.5 times that load, sank only 1 mm in 5 days when the temperature was between -6 and -12 °C. Somewhat similar observations have been reported by T. Andrews. 212

If the force upon the object is inclined to the surface, the ice rises in a hillock in front of the object (Bianconi ²¹³). In all these cases, large blocks of ice were used, and the pressure was probably normal to the surface of freezing, and therefore, parallel to the optic axis, though there is in the

^{*} The logarithmic expression given in their paper seems to be affected by a typographic error, as it requires this b to be negative, which conflicts with their computed values.

²¹² Andrews, T., Proc. Roy. Soc. London, 40, 544-549 (1886).

²¹⁸ Bianconi, J. J., Compt. rend., 82, 1193-1194 (1876).

papers nothing to indicate the direction. Andrews described his data for the rates of penetration at various temperatures as measures of the hardness at these temperatures, and as such they have been quoted, though the property they measure differs from all those commonly classed under that term.

J. T. Bottomley's experiments, 214 in which a loaded block of ice passed through a horizontal sheet of wire gauze, and a loaded wire cut through a block of ice, without in either case permanently damaging the block, are illustrations of both penetration and regelation (p. 412). Bottomley explained them correctly on the theoretical considerations of J. Thomson. 188 The pressure of the wire causes the ice to melt, chilling the wire, the water, and the contiguous ice; the water flows around the wire, thus becoming relieved of stress, and freezes; the heat liberated by the freezing warms the wire, and that the adjacent ice, replacing the heat abstracted by the previous melting, thus preparing the way for a repetition of the process. As shown by Bottomley, the process becomes exceedingly slow—evanescent —if the cold water is drained off before freezing, or if the wire be replaced by a cord. In the first case, the wire is deprived of the heat liberated by the refreezing of the water, and therefore it and the adjacent ice soon become chilled to the temperature corresponding to the melting point of ice under the existing stress; then the melting ceases except as heat is conducted along the wire from the surroundings. In the second case, the cord, a poor conductor of heat, is warmed by the freezing water only on its upper side, and its lower side and the adjacent ice soon become so cool that melting ceases. G. S. Turpin and A. W. Warrington ²¹⁸ repeated Bottomley's experiments, arriving at the same conclusions. They explain Pfaff's observations at temperatures above -1 °C in a similar manner.

See also H. Hess, 108, pp. 14-18; 216

Flowing.—An aggregation of irregular blocks of ice may be welded together by pressure into an apparently uniform mass, and ice may be made to flow through small openings and tubes.

The phenomena considered by J. Thomson (p. 430), i.e., fusion under stress, flow of the melt, and regelation, are contributing factors in most, if not in all, and the controlling factors in many, of the laboratory experiments that demonstrate the flowing of ice (see also J. Thomson 144). But there are other factors that need to be considered, which may in extreme cases become of prime importance, especially when the mass of ice is great and its temperature is low.

Using crystals of NaCl in their saturated aqueous solution, J. Thomson ²¹⁷ demonstrated experimentally the welding and molding of crystals by stresses borne by them but not by the adjacent (saturated) liquid.

O. D. von Engeln 218 has reported the following observations on the

²¹⁴ Bottomley, J. T., Nature, 5, 185 (1872). ²¹⁸ Turpin, G. S., and Warrington, A. W., Phil. Mag. (5), 18, 120-123 (1884). ²¹⁰ Hess, H., Ann. d. Physik (4), 36, 449-492 (1911). ²¹⁷ Thomson, J., Proc. Roy. Soc. (London), 11, 473-481 (1861). ²¹⁸ von Engeln, O. D., Am. J. Sci. (4), 40, 449-473 (1915).

welding of ice by pressure. Into a copper cylinder 4 in. in diameter and 12 in. high, with walls 1/16 in. thick, he placed a rough-hewn plug of pondice with the component crystals parallel to the length of the cylinder. The space between the plug and the cylinder was filled with water, and the water was frozen, thus filling the cylinder with solid ice. A longitudinal pressure of 500 lb*/in², carried in part by the copper cylinder, was applied for several hours; the pressure was then increased to 720 lb*/in², and was maintained over night; the next morning it was increased to 750 and at 4 P.M. to 1400 lb*/in², which was left on for 36 hrs. Then, at 20 °F $(-6.7 \,^{\circ}\text{C})$, the highest temperature reached during the test, the lowest being -4 °F (-20 °C), the pressure was relieved, the cylinder was gently warmed, and the core of ice was slipped out. There had been no distortion of the metal cylinder. The core of ice was "of crystal clearness and homogeneous, showing no line of separation to mark the juncture of the roughhewn prism of pond ice and the water frozen around it. The most striking result, however, was the fact that the ice mass had been completely recrystallized. The original pond-ice core was inserted with principal axes parallel to the pressure direction, the new crystals extended across the cylinder with their principal (and longer) axes at right angles to the pressure direction." They extended "straight across the cylinder instead of radially inward as might have been expected by analogy to the structure of cakes of can-frozen artificial ice."

The crystals were elongated and their terminals were wedge-shaped, thus contrasting sharply with glacier grains. He thought that the difference was due "to the fact that the conditions of our experiment permitted of no movement in the ice mass involved." In other experiments, differing from the preceding principally in the use of a softer metal cylinder, of cracked ice or snow with the crevices sometimes filled with water and frozen, sometimes not, and of a piston fitting into the cylinder so that the entire pressure was borne by the ice, the cylinders were distorted, allowing portions of the ice mass to move. In these experiments also there was a complete recrystallization, but the structure was granular, and the grains were "variously oriented crystallographically." (Such variously oriented grains had been previously reported by A. v. Obermayer.²²⁸)

Whence he concluded "that a granular ice can be developed from snow by pressure with accompanying movement and at air temperatures eliminating the possibility of pressure melting and regelation." The pressures required are near those at which a cube of ice yields when not supported laterally.

The flowing of ice from a large mass through a small opening or tube is merely a special aspect of the penetration experiments of Pfaff,²⁰⁶ Bianconi,²¹⁸ and Andrews,²¹² and is illustrated on a grand scale by the flowing of a glacier through a valley of varying width.

The early tendency to attribute such flowing solely to the lowering of the melting-point by pressure, and the subsequent regelation when the

pressure was reduced, led to a vaguely held idea that the flow is actually that of a liquid, the portion of the ice under pressure being actually lique-This was thoroughly disproved by R. W. Wood 222 who showed by direct experiments that leaden balls embedded in the upper portion of a block of ice remain there even when the block is subjected to a continuously increasing pressure up to 7, 12, and finally 40 tons* per sq. in.;† at the highest pressure, water squirted through the pores of the iron cylinder containing the ice. He also showed that general liquefaction should not have been expected, for, when ice is initially in equilibrium, the application of pressure causes thawing throughout the entire volume of the ice. That abstracts heat, and proceeds only until the entire mass has been reduced to the melting point that corresponds to the applied pressure. This reduced temperature will facilitate the flow of heat from the surroundings into the ice, and will thus facilitate thermal melting, which will be added to that caused by the pressure. All the water so formed is subjected to the pressure, and will escape through all available channels, leaving the bulk of the ice almost as solid as before the pressure was applied. Not only is the resulting water distributed throughout the entire volume of the ice, but the fraction of the ice melted by the direct effect of the pressure is actually small. For example, if the initial temperature of the ice were 0 °C and no heat were allowed either to enter or to leave it, then under a pressure of 7 short-tons*/sq. in. the temperature would fall to -8.5 °C and less than 6 per cent of the original ice would melt; at 15 short-tons*/sq. in. the temperature would be -22 °C, less than 1/6 of the ice would be melted. and the system would be at the triple point (water, ice-I, ice-III); a further increase in pressure would cause freezing and a rise in temperature (see Sections 92 and 93). Some years after this work of Wood's, Sir James Dewar ²¹⁹ performed the same experiment with lead shot embedded in ice, and obtained the same result. They did not sink at all, but were very irregularly distorted. He worked at -80 °C and used a pressure of 100 tons*/sq. in.± (15,000 atm.), believing that the melting-point would by that pressure be reduced to -80 °C. We now know (Sections 92 and 93) that at that temperature and pressure ice exists in the form of ice-VI. not of ice-I, nor of water. (See also Section 93, Fig. 13.)

Wood was interested in adiabatic melting, in distinguishing between "thermo-molten" and "pressure-molten" ice, and approximately obtained that condition, there being no way in which the apparatus could obtain heat except from the surrounding air. He comments on its intense cooling. It is probable that the temperature dropped approximately as has been indicated, and that the fraction of the ice that was actually molten was, as he stated and as the position of the balls indicated, very small.

[†] These seem to be the short tons (2000 lbs*).

[‡]Long ton of 2240 lbs.*

¹⁰⁰ Dewar, Sir James, Chem. News, 91, 216-219 (1905) ← Proc. Roy. Inst. Gt. Brit., 17, 418-426 (1903).

On the other hand, Dewar endeavored to keep the ice at -80 °C by embedding that part of his apparatus in solid CO₂; at that temperature, as we now know, the liquid phase does not exist at any pressure. There is no reason for expecting the shot to sink.

In contrast to Wood and Dewar, A. Mousson²²⁰ had previously performed a similar experiment under approximately isothermal conditions and at such a temperature (-18 to -21 °C) that, as we now know, the liquid phase does exist at certain pressures lower than the maximum (ca. 13,000 atm) reached by him. Under such isothermal conditions, the amount of melting is not limited in the manner considered by Wood, and it is to be expected that during a portion of the process the entire mass of ice will be molten, and that the metal object (in this case a rod of copper) will then fall to the bottom. That he observed. By far the greater portion of the ice was, of course, melted by heat drawn from the surrounding freezing mixture, the true pressure-melting being small; the pressure conditioned, rather than caused, the complete melting. Similar experiments have been reported by J. B. Boussingault.²²¹

If an outlet is provided for the molten ice, it will flow toward it, and will progressively freeze as the pressure on it decreases, the fraction that freezes depending upon the conditions.

Thus there arises a spurious flowing of the ice, in which ice is melted at one point, and the resulting water flows to another, where it is refrozen. Such spurious flowing is always to be expected, but the line of advance of the ice so formed—the birthplace of that ice—is limited to those surfaces at which there is a steep gradient of pressure.

In addition to this, there is a true flowing in which a solid mass of ice undergoes changes in shape. This is shown by the manner in which embedded bodies and superficial landmarks (stakes, rocks, etc.) are carried along by glaciers, and by the flow of ice, under pressure, through contracting conical nozzles, and possibly by the corresponding flow through small apertures.

R. W. Wood ²²² found that at a pressure of 3 tons*/sq. in. ice began to flow through a lateral hole (1/12 in. in diameter) in his compression chamber. It flowed slowly and steadily as a clear cylinder of ice, which broke off when 6 or 8 inches long. At 4.5 tons*/sq. in., "it seemed fairly to spurt from the orifice"; it flowed irregularly, sticking for a second or two and then yielding suddenly. The temperature was about 0 °C.

Sir James Dewar ²¹⁹ has stated that near 0 °C ice can easily be extruded in the form of a wire of clear, transparent ice; at -80 °C such flow still occurred, but "the ice wire was now made up of what looked like a set of disc-like scales"; at a temperature near that of liquid air, "no pressure the

²⁸⁰ Mousson, A., Ann. d. Physik (Pogg.), 105, 161-174 (1858).
280 Boussingault, J. B., Compt. rend., 73, 77-79 (1871) ← Ann. de chim. et phys. (4), 26, 544-547 (1872) → Ann. d. Physik (Pogg.), 144, 326-329 (1871).
282 Wood, R. W., Am. J. Sci. (3), 41, 30-33 (1891).

apparatus would stand caused any flow, but only intermittent explosive ejections."

A similar experiment in which ice was forced to flow through a lateral tube 3/4 in. (1.9 cm) in diameter has been reported by O. D. von Engeln. 218 At a temperature of 22 °F (-5.5 °C), cracked ice was put in a pressure cylinder and by means of a closely fitting piston, a pressure of 3400 lb*/in² (239 kg*/cm²) was applied at 5 P.M. During the night the temperature fell to 4 °F (-15.5 °C), but rose to 14 °F (-10 °C) by 10: 30 A.M., at which time the pressure had fallen to 3000 lb*/in² (211 kg*/cm²) and the extruded core of solid ice was 3 in. (7.6 cm) long; at 4:30 P.M. the core was $5\frac{1}{2}$ in. (14 cm) long, and the temperature was 29 °F (-1.7 °C). The temperature remained approximately that until 10 A.M., when the core was 14 in. (35.5 cm) long. The extruded rod was "perfectly clear, glassy and compact," even at the lowest temperature; the individual crystals were variously oriented; "shear lines and breccia bands could be identified but there were some apparently real crystal boundaries."

In 1902, H. Hess ²²³ forced ice from a cylinder through an attached conical nozzle, and measured the rate of advance of the piston acting upon the ice. In those experiments, essentially the entire flow was through the nozzle. He found there was no flow unless the pressure exceeded a certain value (p_{\min}) depending upon the temperature and upon the ratio of the area of the emerging cylinder of ice to that of the initial cylinder. At 0 °C, he found the following values for (ratio, p_{\min}): (1/9, 345), (1/6.3, 230), (1/3.1, 100), and (1/1.67, 30), the products of ratio times p_{\min} being 38, 36, 32, and 18. The values of (t, p_{\min}) for the fixed ratio 1/6.3 were (0, 230), (-3 to -5, 250), and (-10, 270). In both cases the unit of p is 1 kg*/cm².

When the pressure is kept constant, the rate of flow increases progressively; and after a rate of flow has been established, a reduced pressure will maintain it. This is explained by H. Hess ²²⁴ as due to the lubricating action of the water produced by the melting caused by the pressure.

Phenomena attending the flow of ice through apertures, and the flow of glaciers, have been studied and discussed by A. v. Obermayer ²²⁸ also.

A type of experiment that is intermediate between the simple experiments on penetration and those on the flow of ice through conical nozzles is that in which the ice is enclosed in a cylinder having a diameter greater than that of the piston to be forced into the ice. Under such conditions the piston can advance only as the ice is compressed or as an equivalent volume of ice (or water) is transferred through the gap between the piston and the walls of the cylinder. An interesting experiment in which the gap is wide is described by Hess.²²⁴

In such experiments, G. Tammann,²²⁹ employing small gaps, found that, for any given temperature and pressure, the rate of advance of the

²²³ Hess, H., Ann. d. Physik (4), 8, 405-431 (1902). See also, "Die Gletscher," pp. 28-31, and Ann. d. Physik (4), 36, 449-492 (1911).

²²⁴ Hess, H., Ann. d. Physik (4) 36, 449-492 (1911).

piston is normal unless the pressure is at least essentially as great as that (p_m) at which ice melts at the given temperature. An increase in temperature or in pressure increases the rate; and as the gradually increased pressure passes through p_m , the rate changes abruptly from a small value to a very much greater one. The rates observed for a given apparatus are shown in Table 189. In a later paper ²¹⁰ he represents these data by the formula $\log_{10}r = k(p-p_0)$, where r is the rate of advance of the piston, p is the applied pressure, and k and p_0 are constants for a given temperature and apparatus. Actually, the choice of the values to be assigned to k and to p_0 involves quite an element of judgment, and those chosen satisfy only those observations for which the tabulated value of r exceeds unity. Values of k and p_0 are given at the bottom of Table 189. Similar experiments have been described by N. Slatowratsky and G. Tammann. ²²⁵

Table 189.—Flow of Ice through an Annular Gap 229

r= rate of advance of the piston into a cylinder of slightly greater diameter; pressure on the piston =p kg*/cm²; temperature =t °C. For a given p and t, r is normal if $p \ge p_n$; at $p=p_t$ the rate was too great for measurement; p_m is the hydrostatic pressure at which ice-I melts at t °C. G. Tammann and W. Salge ²¹⁰ represent these data by the formula $\log_{10}r=k(p-p_0)$, where k and p_0 have the values here given. (Sec text.)

	Unit of $p=1 \text{ kg*}/$	cm ² ; of $r = 0.0004538$ c	$em/min = 4.538 \ \mu/min$	•
$t\rightarrow$	-5.7	- 10.7	-15.7	-21.7
100	0.9	0.03		-
200	4.1	0.3	-	
300	11.8	2.00	0.1	-
400	22.5	4.1	0.3	0.15
500	49.5	8.3	1.5	0.3
600	95.0	19.0	5.1	0.5
700		34	12.6	2.5
800		60	2.0	7.0
900		101		13.5
1000		170	*********	20.5
1100	-		***************************************	30
1200				53
1300				65
p_n	642	1116	1729	2000
Ps	665	1130	1787	2100
p_m	678	1225	1681	2170
$10^4 k$	35	29	34	20
Þο	10	190	390	360

In reference to the flowing of glaciers, R. M. Deeley and P. H. Parr²²⁶ remark: "We have seen that glacier ice consists of crystal granules which not only shear freely along planes at right angles to the optic axis, but also undergo changes at their bounding surfaces which enable the mass to suffer

⁹²⁵ Slatowratsky, N., and Tammann, G., Z. physik. Chem., 53, 341-348 (1905).

²²⁶ Deeley, R. M., and Parr, P. H., Phil. Mag. (6), 26, 85-111 (1913).

continuous distortion under stress. The ability of glacier ice to spread out into piedmonts whose upper surfaces are very nearly level also shows that such shear may take place under very small stresses."

M. Matsuyama ^{210a} has expressed the opinion that in the distortion of ice composed of parallel crystals the surfaces between adjacent crystals play a more important role than the gliding planes perpendicular to the optic axis.

The hillock formed around an object forced into a block of ice in simple experiments on penetration, that formed ahead of a loaded object pushed along a surface of ice (Bianconi ²¹³), and the spreading that sometimes occurs at the loaded section when a horizontal bar of ice supported at its ends is loaded in the middle, are all special cases of flowing. The last, the spreading of the bar, occurs only when the optic axis is transverse to the length of the bar, and the load is at least of the order of 12 kg*/cm² (half the crushing load). See Hess.²²⁷

Recovery from stress.—When ice is relieved from stress, it partially returns to its unstressed form and size, provided that the stress has been neither excessive nor too long-continued. Part of the recovery is immediate, and part is progressive. It is the latter, the so-called elastic aftereffect, and especially as it relates to nonuniform stress, that forms the subject of this section. This progressive recovery has been noticed by many.

K. R. Koch 230 has given a few data showing the magnitude and the slowness of the recovery, which indicate that the lower the temperature, the greater is the amount of the progressive recovery, and the longer it takes. For an experiment at -12.5 to -15 °C he records, in arbitrary units, immediate recovery = 12.5, total recovery in 25 min 25 sec = 48.0, followed by an additional recovery of 10 in the next 10 hr 11 min. The total recovery was over 4.6 times the immediate recovery, and required hours.

In another experiment at -1.5 °C he records: Immediate recovery = 30, total in 5 min 15 sec = 41.8, no change in the next minute. Here, the total was only 1.4 times the immediate, and required only 5 min.

In his study of the bending of bars, J. C. McConnel ²³¹ gave special attention to recovery from strain. He wrote: "In several cases after a heavy weight was removed, a slight gradual unbending of the bar took place. At first I thought this a mere consequence of the irregular elastic strains on the bar, the parts most severely strained gradually bending back the rest. But the magnitude of the recovery seems, on closer examination, to put this explanation out of the question, and I have now little doubt that it is a true molecular effect.... I conclude, then, that we have to deal with

²²⁷ Hess, H., "Die Gletscher," p. 21; Ann. d. Physik (4), 8, 405-431 (1902).

²⁰⁸ v. Obermayer, A., Sitzb. Akad. Wiss. Wien [2a], 113, 511-566 (1904).

²²⁰ Tammann, G., Ann. d. Physik (4), 7, 198-224 (1902). 220 Koch, K. R., Ann. d. Physik (Wied.), 25, 438-450 (1885).

²⁸¹ McConnel, J. C., Proc. Roy. Soc. (London), 49, 323-343 (1891).

a real tendency of the forcibly displaced sliding layers to slide back. The rate of recovery, rapid at first, soon falls off."

M. Matsuyama ^{210a} has given certain data for the recovery of rods of ice from torsion about the axis of the rod, but information concerning the time allowed for that recovery seems to be lacking.

Brittleness.—E. Reusch ^{209a} observed that brilliant cracks, like those produced in glass by means of a diamond, can be produced in ice by pressing upon it with a convex knife-blade; and that these cracks can be formed even when the ice is in a warm room, and consequently, is covered with a layer of water. This indicates that ice is brittle even when near its melting point. G. Tammann and W. Muller ²³² have stated that at 0 °C ice is as brittle as is rock salt when 700 °C below its own melting point.

Nevertheless, ice yields progressively to the action of differential stresses even when below its melting point. It is both plastic and brittle.

In speaking of the work of E. Brown,²³³ H. T. Barnes says that ice splinters considerably when sawed at temperatures near $0 \,^{\circ}\text{F} \, (-17.8 \,^{\circ}\text{C})$, but it can be sawed at $30 \,^{\circ}\text{F} \, (-1.1 \,^{\circ}\text{C})$ with comparatively little difficulty.²³⁴

Quantitative Treatment.

Young's modulus.—The determination of the value of Young's modulus of elasticity (E) of ice by the usual static methods is rendered very difficult by the progressive yielding of ice to stress, and by its partial progressive recovery (elastic after-effect) when the stress is removed. Indeed, it is questionable whether significantly useful values can be obtained by such methods (see Boyle and Sproule, 1931). Nevertheless, most of the values commonly cited have been so derived, and are given in the last two sections of Table 190, as a matter of historical interest.

On the other hand, the dynamic methods based upon the velocity of propagation of high-frequency vibrations lead to values of E that are unaffected by the progressive yielding (see Boyle and Sproule, 1931). They alone deserve serious consideration.

Values obtained by the static method exhibit wide variations, and have been interpreted as indicating that the value of E depends upon the angle between the stress and the optic axis. This has not been borne out by the results obtained by the dynamic method, which indicate that values of E found for specimens that are nominally identical may differ by some 10 to 15 per cent, and that the observed variations with the orientation of the optic axis are of about the same magnitude. That is, there is no certain dependence of E upon the orientation of the crystal.

The observations of Trowbridge and McRae (1885), and some of those of Hess (1902, 1904) indicate that, when the apparent value of E is derived from the bending of a loaded horizontal bar, the planes of freezing being

²⁸² Tammann, G., and Muller, W., Z. anorg. allyem. Chem., 224, 194-212 (1935).

²⁸⁸ Brown, E., Rep. Joint Board Eng. for St. Lawrence River.

²⁸⁴ Barnes, H. T., "Ice Engineering," p. 224, 1928.

Table 190.—Young's Modulus of Ice

The most accurate values are those of Boyle and Sproule, of which the individual determinations in any given case lie within a range of 3.5 per cent after correction has been made for differences in the temperatures; different specimens, nominally identical, occasionally differed by 15 per cent. They used longitudinal vibrations of frequencies between 7 and 13 kilocycles/sec.

Values obtained by the static method are unreliable (see text) and mainly of historical interest.

 θ = angle between the length of the specimen and the normal to the surface of freezing; E = Young's modulus; l = length of the specimen, w = width, τ = thickness in direction of application of the load; Op. Ax. = optical axis.

Unit of $E = 1 \text{ kg}^{+}/\text{mm}^{2} = 1424 \text{ lb}^{+}/\text{in}^{2} = 98.1 \text{ megadyne/cm}^{2}$. Temp. = $t \, ^{\circ}\text{C}$

I. Dynamic method. Longitudinal vibrations except as noted.

	Boyle and Sproulea; $\theta = 0^{\circ}$					yle and Spro	ules: $t = -2$	6 °C
	-9	-10	-30	-35	θ	0°.	45°	90°
\boldsymbol{E}	947	967	1040	1110	E	970	900	9906
E_c	954	960	1060	1090	\boldsymbol{E}			9450
E	$_{c} \equiv 909(1 \cdot$	-0.00558t)		E_c	1040		

Miscellaneous values; $\theta = 90^{\circ}$.

Vibration		Longit	udinal		T	rans.	3
t	- 26	- 26	-6	-4	-7	0(?)	?
E	990	9450	880	960	884	236	710^d
Ref.	BS	BS	TMcR	RS	K85	R	\mathbf{Ko}

II. Static method. Bending of loaded horizontal bars.

Sheet ice.

Unless another direction is indicated, the load is applied perpendicular to the surface of freezing, if $\theta = 90^{\circ}$. (From the data given on p. 611 of Matsuyama's article it may be seen that the unit he used and called the "c.g.s." unit is actually 1 gram-weight per cm², not 1 dyne/cm².)

Op. Ax.	111	r E	lw	Ref.a	· · · · · · ·	0°	E -90°	Ref.
0 to -1	182		383	H	-5.4		642	K85
$-2 \text{ to } -5 \\ -1 \text{ to } -5$	59	254	418	H H	-5 to -7 -6.5 to -7.8	609	860 622	TMcR K13
(3)	67	194	336	H	-6.5 to -7.8		656 ^f	K13
-3.5	185	60	92	Ma	-9	696	696	K85
					(3)	1120	958	K14
					(?)		950	Mo
					(3)		500	В

Granular ice, natural and compressed.

	R	Type of ice	Ref.a
-1 to -3	285	Large grains	H
0 to -3	226	Small grains	Ĥ
-0.5	300	Mixed sand and water, frozen	H
- 2	150	Snow compacted by 33 atm	H
0 to -0.2	49	Snow compacted by 20 atm	H
0 to -3	280	Crystals randomly oriented	H
-6(?)	620°	Crystals disordered	TMcR
3	190°	Crystals disordered ^h	F

Table 190—(Continued)

 b By a similar method, M. Ewing, A. P. Crary, and A. M. Thorne, Jr., 205 found $E=935~{\rm kg^{*}/cm^{2}}$ at t between -5 and $-15~{\rm C}$; and obtained the same value for ice artificially frozen in a vertical tube as for a rod cut with its length parallel to the surface of freezing.

*Length perpendicular to the preceding, but both parallel to the surface of

freezing.

d Isotropic lake ice.

Direction of application of the load is not stated.

Direction of application of the load is parallel to the surface of freezing.

⁹ By transverse vibrations.

*The ice was frozen in a metal tube; the orientation of the crystals is neither ordered nor perfectly at random.

'Computed from the observed elongation under tension within his estimate of the elastic limit (load $< 0.51 \text{ kg*/cm}^2$).

horizontal (optic axis vertical), as the load increases the apparent value of E decreases to a minimum, and then increases as the breaking load is approached. This variation is, however, small as compared with that of the apparent E from specimen to specimen.

Data reported by O. Fabian (1877) indicate that ice is almost perfectly elastic for tensions not exceeding 0.5 kg*/cm², but it should be noticed (Table 190) that the value he obtains for E under such conditions is very low.

Poisson's ratio.—As computed from the observed velocity of longitudinal waves, Poisson's ratio for ice is 0.365 ± 0.007 , and is the same for ice frozen in a vertical tube as for a rod cut with its length parallel to the surface of freezing,²³⁵ and from the horizontal velocity of waves in an isotropic ice-sheet, R. Köhler ²³⁶ derived the value 0.30. From static observations, B. Weinberg ²³⁷ had derived the value 0.38 ± 0.49 ; the direction of extension is not clearly indicated.

Rigidity.—The rigidity of ice has been derived by B. Weinberg,^{237, 238} K. R. Koch,²³⁹ M. Matsuyama,^{210a} and C. D. Hargis,²⁴⁰ from observations on the torsion of bars, Weinberg deducing it from the apparent viscosity by means of an extension of Maxwell's theory (see *Plasticity*, p. 451),

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<sup>235</sup> Ewing, M., Crary, A. P., and Thorne, A. M., Jr., Physics, 5, 165-168 (1934).
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²⁸⁶ Köhler, R., Z. Gcophys., 5, 314-316 (1929).

²⁸⁷ Weinberg, B., Z. Gletscherkunde, 1, 321-347 (1907).

²⁸⁸ Weinberg, B., Ann. d. Physik (4), 22, 321-332 (1907).

²⁸⁹ Koch, K. R., Idem, 45, 237-258 (1914).

²⁴⁰ Hargis, C. D., Phys. Rev. (2), 19, 526, 527 (1922).

and the others from the twist produced by a known torque, care being taken to eliminate the effect of plastic yielding and of the elastic after-effect. Weinberg calculated also the relaxation time (τ) , and the strain (λ) corresponding to the elastic limit. There appears to be an error in the equation used by him in those computations (see p. 453), but the size of the error so introduced into the computed values is not known. From the numerical data on p. 611 of Matsuyama's paper it may be shown that the unit which he designates by "c.g.s." is actually 1 gram-weight per (cm²-radian), although in the table on p. 615 he compares such values with others which are actually expressed in terms of 1 cgs unit = 1 dyne/(cm²-radian).

The dynamic determination—from high-frequency torsional vibration of rods—by M. Ewing, A. P. Crary, and A. M. Thorne, Ir.²³⁵ yields a value that is markedly higher than those obtained by the static methods, but which is probably to be preferred, though as yet unsupported.

Table 191.—Rigidity of Ice

N = modulus of rigidity; $\theta = \text{angle}$ between length of cylinder and optic axis; λ = shear corresponding to the elastic limit; τ = relaxation time (p. 452); E = Young's modulus; $t \circ C = temperature$.

N is determined from the twisting of cylinders by torques about the axis of figure. Weinberg obtained $N = 1.0 (1 - 0.13t) 10^{10} \text{dyne/(cm}^2 \cdot \text{radian)}$ when $\theta = 0$, and N = 0.8 (1 - 0.65t) for glacier ice. Matsuyama reports for $\theta = 90^{\circ}$, $N = 1.16 (1 - 0.080t - 0.0017t^2) 10^9 dyne/(cm²-radian). GI =$ glacier ice.

Unit of N=1 kmegadyne/(cm²-radian) = 1019 kg*/(cm²-radian); of $\tau=1$ sec, of $\lambda=1$ microradian=0.206"

Preferred value. Dynamic method. ECT.^b

 $N = 91.7 \pm 0.5$; t = -5 to -15 °C. Water frozen slowly in vertical brass tube.

II. Static or slow oscillation method.

<i>0</i> →	0	90	G1	0	Gl	$\frac{0}{r}$		0 8/3	y 90	Ref.
t										**
-	27.2	29.4	-	-	-		-	4.03	3.18	\mathbf{K}
0	10		8	56	10	960	480			W
-5	17	-	34	340	3	1670	720			W
-5		1.6				-				\mathbf{M}
-6	1.8	-					·			\mathbf{M}
	28.2^{d}			-	-					H

At an unstated temperature and for an isotropic ice-sheet, R. Köhler 256 found N = 27.

* References:

Ewing, M., Crary, A. P., and Thome, A. M., Jr. 233 Hargis, C. D. 240 Koch, K. R. 250 Matsuyama, M. 210a Weinberg, B. 238 ECT H K

° In finding λ , N was taken as 16.5.
d Value of θ is unknown; the ice was frozen in a brass tube.

Tensile strength.—The tensile strength of ice may be expected to depend upon the structure of the ice and upon the direction of the line of stress with

reference to the optic axis of the crystal, or crystals, of which the specimen is composed. No data on the tensile strength of single crystals or of ice of known uniform structure have come to the compiler's attention. The recorded values, at unstated temperatures, range from 2.4 to 16 kg*/cm², and Reusch computes 68 kg*/cm² as the maximum tension at the instant of rupture of a centrally loaded rod supported near its ends.†

Working at -8 °C and loading the specimen at the rate of 0.1 kg*/(cm²sec), H. Romanowicz and E. J. M. Honigman ²⁴¹ found in three tests the mean values 16.1, 18.3, and 17.7 kg*/cm², the highest observed value being 24.8, and the lowest 14.8 kg*/cm².

Strength in linear compression.—The values reported for the linear compressive stress required to rupture a block of ice vary from 5 to 125 kg*/cm² (70 to 1800 lb*/in²), the values most frequently found lying near 25 kg*/cm² (360 lb*/in²). This wide variation is in large part due to variations in the structure of the ice, but in part to the technique employed. Barnes,^{234, p. 228} quoting Prof. E. Brown.²³³ says that the observed crushing stress depends upon the rate at which the stress is applied. That rate is seldom reported. The crushing stress may depend also upon the size of the specimen (G. van Diesen, 1871).

For 7-cm cubes loaded at the rate of 3 kg*/(cm²sec), H. Ramanowicz and E. J. M. Honigman ²⁴¹ found in three tests the mean values 40.0, 43.0, and 44.1 kg*/cm², the extreme observed values were 54.4 and 34.0; the ice was formed by freezing water in cubic forms a little larger than the desired finished block.

No data for an isolated crystal have come to the compiler's attention, the data available referring to blocks of natural or of ordinary artificial ice, which consist of multitudes of crystals seldom arranged in more than approximate uniformity. In some cases the average direction of the optic axes was inferred from the direction of the planes of freezing, being assumed to be normal to those planes. For the values obtained in such cases, see Table 192.

Barnes ²⁴² has reported that when blocks of ice are subjected to linear compression they may be heard to crack at approximately half the pressure required to crush them; they then stiffen perceptibly. He was unable to see these cracks. When the line of pressure is normal to the surfaces of freezing "the ice bursts sideways into innumerable long needles, resembling a cake of ice which has all but fallen to pieces in the sun." When the line of pressure was parallel to those surfaces "the block cracked lengthwise and transversely without shattering." The required pressure is somewhat greater in the first case than in the second.

[†] See: Barnes, H. T., Hayward, J. W., and McLeod, N. M., Trans. Roy. Soc. Canada III, (3), 8, 29-49 (1914); Fabian, O., Rep. f. Exper.-Physik (Carl), 12, 397-404 (1876) ← Sitz.-ber. d. Krakauer Akad. Wiss. (Math.-Nat. Kl.) Vol. 3; Finlayson, J. N.²⁴⁸; Hess, H., "Die Gletscher," p. 23; (Canon) Henry Moseley, Phil. Mag. (4), 39 1-8 (1870); Reusch, E., Ann. d. Physik (Poyg.), 121, 573-578 (1864).

 ²⁴¹ Romanowicz, H., and Honigman, E. J. M., Forsch. Gebiete Ingenieurw., 3, 99 (1932).
 ²⁴² Barnes, "Ice Engineering," 1928, p. 220. Trans. Roy. Soc. Canada, III (3), 8, 19-22 (1914).

Von Engeln reported that when a pressure approaching the crushing value was released, the ice frequently cracked, and if the release was rapid it actually broke apart, showing that it retained its elasticity. But if the pressure was maintained near the crushing value the ice yielded by flow without breaking.

Under given conditions, the strength increases as the temperature is reduced.

Taking 400 lb*/in² as the crushing stress, Barnes, Hayward, and McLeod computed the following values for the greatest possible thrusts per transverse linear foot (30.5 cm) when the ice has the thicknesses indicated.

Thickness	{ 6	8	10	12 in.
	15.2	20.3	25.4	30.5 cm
Thrust	28800	38100	48000	57600 lb*/ft
	42900	57200	71500	85700 kg*/m

Other references:†

Table 192.—Strength of Ice in Linear Compression

The following data refer to ice of which the structure is believed to be uniform, the optic axes of the crystals being perpendicular to the surface of the water on which the ice was formed. Stress || (\perp) indicates that the compression is parallel (perpendicular) to the optic axis.

Unit of Strength = 1 kg*/cm ² = 14.24 lb*/ft ² = 0.981 megadyne/cm ² .	Temp. = $t ^{\circ}$ C
---	-------------------------

Stress→	Stre	ngth —	Source	Observer ^a
0	26	25	River	Barnes
-2.2		21	River	Brown
- 10		49	River	Brown
-16.6		62	River	Brown
-11.7	124	72	River	Finlayson
-7	70	25	Pond	von Engeln

^a Barnes, H. T.²⁴²; Brown, E.²³³, ²³⁴, pp. ²²³–²²⁸; von Engeln, O. D., Am. J. Sci. (4), 40, 449-473 (1915); Finlayson, J. N.²⁴³

Shearing strength.—J. N. Finlayson 243 reported that at temperatures above 20 °F (-6.7 °C) the shearing load had to be applied in his experiments "quite rapidly in order to secure satisfactory results."

"In cases where the specimens were sheared in the direction parallel to the optical axes of the crystals, beautiful conchoidal fractures were frequently obtained, indicating that the specimens had sheared along the walls of crystals."

The average of his values for the shearing strength was 114 lb*/in² (8.0 kg*/cm²) perpendicular to the optic axis, and 98 lb*/in² (6.9 kg*/cm²) parallel to that axis. Observations extending from -10 °F to +30 °F

[†] Barnes, H. T., Hayward, J. W., and McLeod, N. M., Trans. Roy. Soc. Canada, III (3), 8, 29-49 (1914); van Diesen, G., Vers. en Med. K. Akad. Wet., Amsterdam (2), 5, 325-331 (1871); von Engeln, O. D., Am. J. Sci. (4), 40, 449-473 (1915); Finlayson, J. N. 563; Hess, H., "Die Gletscher"; Moseley, (Canon) Henry, Phil. Mag. (4), 39, 18 (1870); Anon., Z. ges. Kältelnd., 33, 84-85 (1926), a few values derived from observations made at the government's testing bureau at Copenhagen are cited anonymously.

248 Finlayson, J. N., Canadian Engineer, 53, 101-103 (1927).

 $(-23.3 \, ^{\circ}\text{C})$ to $-1.1 \, ^{\circ}\text{C})$ gave no indication that the strength varies with the temperature. Individual determinations differ widely. A few tests of artificial ice indicated that its shearing strength is about 80 per cent of that of river ice.

"No marked elastic limit was noticeable before the specimens sheared off; but there was evidence of a slow realignment of crystals under pressure, as the load was found to fall off if the head of the testing machine were brought to rest during the test."

These conclusions of Finlayson are quoted by H. T. Barnes.²³⁴, p.217 Weinberg's data (Table 191) indicate that the elastic limit in shear is certainly less than 2 kg*/cm² and probably not over 1 kg*/cm², the limiting shear being a few seconds of arc (0.6" to 11").

Canon Henry Moseley ²⁴⁴ found no detectable shearing of a cylinder of ice 1.5 inches in diameter when the shearing force was 112.5 lbs*, but an appreciable shearing when the load was 121 lbs*; the temperature was stated to have been below freezing. The shearing apparatus consisted of two boards of hard wood held together by guides, and sliding one over the other; a 1.5-inch hole, to take the ice, was bored through each board. With this apparatus he found that a load of 208 lbs* caused shearing at the rate of 0.016 in/min; working in air at 74 to 75 °F he found that 200 lbs* gave a rate of 0.025 in/min for solid, natural ice, and 0.036 in/min for regelated ice formed by hammering cracked ice into the hole in the shearing apparatus. His conclusion was that the shearing strength of such compacted ice is about 75 lb*/in² (=5.3 kg*/cm²)²⁴⁶.

Hardness.—Three kinds of hardness are commonly recognized. One indicates the resistance to abrasion; a second, the resistance to denting under the action of a dead load; and a third, the height of rebound of a specified object dropped in a specified manner.

The first is generally used in the description of minerals, and is indicated most frequently in terms of a scale defined arbitrarily by a specified set of minerals. On Moh's scale, the hardness of ice is generally given as 1.5; that is, its resistance to abrasion lies about midway between that of talc and that of gypsum.^{234, p. 18; 247, 248} E. S. Dana ²⁴⁹ does not state the hardness of ice, but on the basis of several references, it is believed that either earlier editions of this work or J. D. Dana ²⁵⁰ gave the value of 1.5.

The second and the third kinds of hardness are commonly used in the description of metals, and are specified, respectively, by what are known as the Brinell hardness number and the Shore scleroscope hardness. No data for either of these have been found for ice, although there is no obvious

²⁴⁴ Moseley, Canon Henry, Phil. Mag. (4), 42, 138-149 (1871).

²⁴⁵ Moseley, Canon Henry, Phil. Mag. (4), 39, 1-8 (1870).

²⁴⁶ Moseley, Canon Henry, Proc. Roy. Soc. (London), 17, 202-208 (1869).

²⁴⁷ Van Horn, F. R., "General and Special Mineralogy," p. 458, published by the author, Cleveland, 1903.

Bayley, W. S., "Descriptive Mineralogy," p. 147, New York, 1917.

²⁴⁹ Dana, E. S., "A Text-Book of Mineralogy," 3rd ed., revised and enlarged by W. E. Ford, New York, 1922.

²⁵⁰ Dana, J. D., "A Manual of Mineralogy."

reason for anticipating any serious difficulty in determining the Shore scleroscope hardness of ice at any temperature. It would, without doubt, vary with the structure of the ice, and, for homogeneous ice, it would depend upon the direction of the crystallographic axes of the individual crystals. The Brinell hardness number for ice would have no significance except at temperatures so low that the rate at which ice yields progressively under the action of a constant load applied to a small area of its surface is negligible; say, at temperatures below $-30\,^{\circ}\text{C}$.

Certain experiments on the rates at which loaded rods and tubes penetrate into ice at given temperatures have been made by several experimenters, including T. Andrews ²⁵¹ who designated his data as measures of hardness. They have been so quoted, e.g., by H. T. Barnes, ^{234, p. 47; 252} although it is evident that they refer to progressive deformability rather than to hardness in any of the senses in which that term is commonly used. In this compilation they have been assigned to the section *Penetration* (p. 436).

Plasticity and viscosity.—Ice is a plastic solid. That is, under the action of small differential stresses it seems to be perfectly elastic, suffering no permanent change in form, but if the stress exceeds a certain small "elastic limit," its deformation continuously increases. Other things being the same, the nearer the temperature is to 0 °C, the more rapidly does the deformation increase. When the stress is a shear of the type produced by an axial twist applied to one end of a cylinder while the other is held fixed, the value of the strain at the elastic limit, as computed by B. Weinberg ^{237, 238} is independent of the temperature. The phenomena are complicated by the effect of the stress upon the melting point of the ice, especially of the impure intercrystallic material. Many early observers concluded that under differential stresses ice yields progressively, however small the stress may be, especially when the temperature is near 0 °C: J. Thomson, ²⁵³ F. Pfaff, ²⁰⁶ J. J. Bianconi, ²¹³ T. Andrews. ²⁵¹

Both the elastic limit and the rate of yielding vary with the structure of the ice, and, when the structure is uniform, with the directions of the stresses with reference to the optic axes of the constituent crystals. They would also be expected to vary with the amount and nature of the impurities contained in the intercrystallic material. It seems that this last has not yet been considered by those who have investigated the plasticity of ice.

Under the titles plasticity, viscosity, and hardness, many observations of the manner in which ice yields to differential stresses have been reported. Most of them, though very interesting, have been made under such conditions or reported in such deficiency of detail as to make quantitative interpretation impossible. They will be found in the earlier portions of this section. Those from which the observers have attempted to derive numerical values for what they call the viscosity will be considered here. But first

 ²⁵¹ Andrews, T., Proc. Roy. Soc. (London), 40, 544-549 (1886).
 ²⁵² Barnes, H. T., "Ice Formation," p. 66, New York, 1906.

²⁵⁸ Thomson, J., Proc. Roy. Soc. (London), 8, 455-458 (1857).

it is necessary to define the terms we shall use. Consider a plastic material bounded on opposite sides by planes that are parallel and distant x one from If one of these planes is kept at rest and the other in motion in its own plane with a velocity v, then each will experience a drag amounting to P units per unit area, such that $(P - p) = \mu v/x$, μ being a property of the solid but independent of the value of P, v, and x. Unless P exceeds b, v is zero. The force required to produce the strain corresponding to the elastic limit is p units per unit area. This is the form of equation demanded by J. Clerk Maxwell's theory of viscosity 254 as extended by T. Schwedoff ²⁵⁵ to include the case in which there is a definite fixed elastic limit different from zero. It may for convenience be regarded as a definition of a plastic solid, and is in effect the definition adopted by E. C. Bingham.²⁵⁶ Whether any specified solid that is commonly described as plastic satisfies this definition is another question, and one that need not detain us. It is merely a matter of definition; if the solid does not behave in accordance with that definition, then it is not purely plastic in the sense in which we shall use the term.

As our defining equation follows from an extension of Maxwell's theory of viscosity, which provides for other phenomena observed in the study of the shearing of ice, it is desirable to recall the essentials of that theory. He regards a viscous substance as consisting of one or more types of molecular aggregate. When any aggregate is strained by a relative motion of adjacent parts of the substance, it gradually breaks up, relaxing the strain, and the parts then form new associations, not necessarily of the same type of aggregation as before. Thus the strain is gradually relieved unless continually renewed by a continuous relative motion imposed from without upon the adjacent parts of the substance. If the rate of relaxation is directly proportional to the strain, then, if left to itself, the strain decreases exponentially with the time, and the time required for it to decrease to e^{-1} (=0.3679) of its value was called by Maxwell the relaxation time, and denoted by T. The viscosity of the substance is equal to the product of the modulus of rigidity multiplied by the relaxation time.

In general, the value of T will differ from one type of aggregate to another. When a substance containing several types of aggregates is strained, those for which T is small will soon relax, throwing additional stress upon those for which T is great, and they in turn and in some measure protect from stress the newly formed aggregates for which T is small. Thus when the distorting stress is maintained constant the effective viscosity will increase with the duration of the stress; at the same time. the velocity with which the distortion increases will decrease.

When the distorting stress is removed, there will be a partial and progressive recovery of the original form. The aggregates that had been

²⁸⁴ Maxwell, J. C., Phil. Mag. (4), 35, 129-145, 185-217 (1868); "Scientific Papers," 2, 26-78 (1890) = Phil. Trans., 157, 49-88 (1866).

285 Schwedoff, T., Jour. de Phys. (2), 8, 341-359 (1889); 9, 34-46 (1890).

²⁵⁰ Bingham, E. C., Bull. Bur. Standards, 13, 309-353 (SP 278) (1916).

strained less than their elastic limit will recover at once, and in so doing will strain other aggregates. Those will yield elastically or viscously, depending upon the amount they are strained, thus introducing new strains; and so on. In the end, the substance will be subjected to permanent internal strains, unless the elastic limit of each type of aggregate is actually that of zero strain, in which case the substance is purely viscous.

If the aggregates do not relax unless the strain exceeds a certain value, the substance is plastic. In general, that limiting strain will differ from aggregate to aggregate, and may have any value from zero (purely viscous) to infinity (perfectly elastic for all stresses). The behavior of the substance under shearing stress will vary accordingly.

Those who have studied ice have called Px/v the viscosity. We shall call it the effective viscosity, and shall denote it by μ_c ; and we shall call the

quantity denoted by μ in the equation $(P - p) = \frac{\mu v}{x}$ the viscosity. This

does not accord entirely with the somewhat confused nomenclature used by those interested in the study of plastic materials used in the arts, but it is logical and is justified by the manner in which the defining equation was derived from Maxwell's picture of the structure of viscous substances. In our notation $\mu_0 = \mu + px/v$.

Although several experimenters have observed that μ_e for ice increases as v becomes small, B. Weinberg 257 appears to be the only one who has attempted to separate the two terms composing μ_e . From a consideration of the progressive yielding of cylinders of ice, each clamped at one end and subjected at the other to a constant torque about the axis of figure, he concluded that his observations can be quite satisfactorily expressed by formula (1). (In the original paper, the negative sign has been omitted from the exponent.)

$$\mu_c = \mu_0 \left(a - \frac{b}{t} \right)^{-t} + \frac{c}{\psi} \tag{1}$$

Here, the temperature is t °C, the rate of shear is ψ radians per sec, and μ_0 is the value of the viscosity at 0 °C and $\psi=\infty$. He does not state how long the stress had lasted when ψ was observed. For river ice, the geometrical axis of the cylinder being parallel to the optic axes of the constituent and parallel crystals, he gives $\mu_0=9.5$ megamegapoises, a=1.12, b=0.54 °C, and c=0.5 megadyne-radian/cm². For glacier ice he gives $\mu_0=3.8$ megamegapoises, a=1.32, b=0.65 °C, and c=0.08 megadyne-radian/cm². From these he computes the modulus of rigidity, the relaxation time, and the greatest shear for which there is no permanent deformation (see Table 191). This maximum shear at 0 °C is 56 microradians for the river ice, and 10 for the glacier ice. These computations are vitiated by an error analogous to that considered by E. Buckingham 258 in his dis-

Weinberg, B., Ann. d. Physik (4), 22, 321-332 (1907). Superseding and extending Idem, 18, 81-91 (1905).

⁹⁸⁸ Buckingham, E., Proc. Amer. Soc. Testing Materials, 21, 1154-1161 (1921).

cussion of an equation used by Bingham to represent the flow through a capillary tube. Values defined by means of formula (1) are given in Table 193.

Table 193.—Viscosity of Ice 257

 $\mu_e = \mu_0 \left(a - \frac{b}{t} \right)^{-t} + \frac{c}{\psi}$; temperature = t °C; rate of shear = ψ radians/scc, corresponding to a difference of v meters/year in the velocities of two planes of slipping that are 100 meters apart. In his paper of 1905 Weinberg ²⁵⁷ gives $\mu_t = (12.44 - 4.02t + 0.277t^2) \times 10^{12}$ poises when the mean value of ψ is about 10^{-8} radian/sec; this formula is probably not so good as the other. Computation by the compiler.

Unit of $\mu = 10^{12}$ poises, of ψ and of v as already indicated; temp. = t °C

I. River ice. Planes of slipping are perpendicular to optic axis. $\mu_0 = 9.5$, a = 1.12, b = 0.54 °C, $c = 5 \times 10^5$ poise radian/sec.

$\psi \rightarrow$	10-8(?)	5×10-9	10-8	10-7	5×10^{-7}	∞
2/→	31.6(?)	15.8	31.6	316ª	631	∞
Z	μ_t					
0	12.4	110	60	14.5	10.5	9.5
-0.1	12.8	112	62	16.5	12.5	11.5
-0.5	14.5	114	64	19.1	15.1	14.1
-1.0	16.7	116	66	20.8	16.8	15.8
-2.0	21.6	118	68	23.3	19.3	18.3
2.0	21.0	110			17.0	
-3.0	27.0	121	71	25.9	21.9	20.9
-4.0	33.0	124	74	28.6	24.6	23.6
-5.0	39.5	126	76	31.5	27.5	26.5
- 7.5	58.2	135	85	40.3	36.3	35.3
- 10.0	80.3	147	97	52.3	48.3	47.3
-12.5	93.0	163	113	67.3	63.9	62.9
-15.0	135	184	134	88.6	84.6	83.6

II. Glacier ice. $\mu_0 = 3.8$, a = 1.32, b = 0.65 °C, $c = 8 \times 10^4$ poise-radian/sec.

$\begin{matrix} \psi \rightarrow \\ \tau \rightarrow \\ t \end{matrix}$	10 ⁻⁹ 3.16	5×10 9 15.8	10 ⁻⁸ 31.6	10 7 316ª	5×10 ⁻⁷ 631	%
o	83.8	19.8	11.8	με	4.0	3.8
-0.1	84.7	20.7	12.7	5.5	4.8	4.7
-0.5	86.1	22.1	14.1	6.9	6.3	6.1
-1.0	87.5	23.5	15.5	8.3	7.6	7.5
-2.0	90.3	26.3	18.3	11.1	10.5	10.3
-3.0	93.8	29.8	21.9	14.6	14.0	13.8
-4.0	98.4	34.4	26.4	19.2	18.5	18.4
-5.0	104.4	40.4	32.4	25.2	24.5	24.4
-7.5	129	65.0	57.0	49.8	49.2	49.0
-10.0	178	114.4	106.4	99.2	98.6	98.4
-12.5	278	214	206	198	198	198
-15.0	475	411	403	396	395	395

[&]quot;At this rate, two planes of slipping that are 10 cm apart will differ in velocity by 36 microns per hour.

In his earlier paper, Weinberg gives formula (2) for the shearing of river ice in a direction perpendicular to the optic axis, and at rates of approximately 0.01 microradian per second.

$$\mu = 12.44 - 5.02t_R + 0.355t_R^2$$
 megamegapoises
= 12.44 - 4.02t + 0.227t² (2)

where t_R is the temperature on the Réaumur scale (not Centigrade scale, as first published; see the 1907 paper). Values computed by means of this formula are given in the second column of the first part of Table 193.

From an extended study of the bending of horizontal rectangular bars of ice, supported near each end, and loaded in the middle, H. Hess 258a had already deduced values for the apparent viscosity. These values were

computed by means of the formula $\mu_e = \frac{lP}{4abv} = \frac{M}{v}$, where $M \equiv \frac{lP}{4ab}$, P =

load, l = length between supports, a = vertical thickness, b = horizontalbreadth, v = velocity of depression of the mid-point of the bar = $l\psi/2$, where ψ radians/sec is the rate of shear; ψ does not exceed a few times 10^{-8} . These values of μ_e are less than a hundredth as great as those found by Weinberg, and vary with the length of time the load has been applied. He reported that there was no detectable change in μ_e in the range 0 to -6.8 °C. For three bars cut from the same sheet of uniform ice he recorded the values given in Table 194. For granular ice he obtained values of the same order of magnitude. Successive bendings in opposite directions produced no change in the value of μ_c , the load being moderate. From his observations, he concluded that, under moderate loads, μ_0 increases with the duration of the load, and after about 5 min the rate of increase is essentially constant; but under loads near the breaking value, μ_e decreases as the duration increases. Such variations may be forecast from the extension of Maxwell's theory. For additional details, reference should be made to the original paper.

Similar observations on the bending of loaded bars of ice had been made 11 years earlier by J. C. McConnel 231 in his very important reconnaissance of the behavior of such loaded bars. He thought it unprofitable to attempt to compute the viscosity from his observations, but R. M. Deeley 250 has made such computations, finding for shears perpendicular to the optic axis values of the same order (1010 poises) as those obtained by Hess. The compiler has been unable to obtain from McConnel's data the actual values published by Deeley. McConnel's observations show that the apparent viscosity for shear parallel to the optic axis is many times (perhaps 100) as great as that for a shear perpendicular to that axis.

From the observations of J. C. McConnel and D. A. Kidd 260 on the progressive elongation of bars of ice subjected to longitudinal traction, R. M. Deeley and P. H. Parr 226 have computed the apparent viscosity,

²⁵⁹a Hess, H., Ann. d. Physik (4), 8, 405-431 (1902); "Die Gletscher" (1904).

²⁰⁰ Deeley, R. M., Proc. Roy. Soc. London (A), 81, 250-259 (1908).
200 McConnel, J. C., and Kidd, D. A., Proc. Roy. Soc. (London), 44, 331-367 (1888).

finding values varying from 9 to 900 megamegapoises, depending upon the temperature, the structure of the ice, and the direction of the shear.

Table 194.—Viscosity of River Ice 258a

Values were derived from the bending of horizontal, rectangular bars supported at the ends and loaded at the middle. P = load; M = bending moment per unit of cross-sectional area = Pl/4ab; $\mu_{\tau} = \text{value}$ of the apparent viscosity as computed from the rate of shear τ sec after the load was applied; l = length between supports; a = vertical thickness; b = horizontal breadth; vertical and horizontal refer to position of bar when loaded for test. All three bars were cut from the same sheet of ice.

Unit of $P = 1$ g*, of $M = 1$ g*·cm ⁻¹ , or	$f \mu =$	1012 poises
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Axis→		Parallel to l			Parallel to a	
<i>P</i> → <i>M</i> →	2000 1350	5000 3400	6000 4000	1000 1600	1500 2350	2000 3100
τ		μτ			μτ	
15	0.065	0.105	0.0055	0.075	0.100	0.080
60	0.175	0.115	0.036	0.075	0.110	0.070
120	0.100	0.130	0.0365	0.075	0.090	0.110
300	0.110	0.160	0.035	0.080	0.120	0.120
1200		0.120				

		Optic axis p	arallel to b	
<i>P</i> → <i>M</i> →	1000 1500	1500 2250	2000 3000	3000 4450
τ		μ	l _T	
15	0.037	0.037	0.024	0.110
60	0.080	0.110	0.060	0.090
120	0.120	0.100	0.100	
300	0.210	0.190	0.170	

In the same paper, Deeley and Parr summarize the more important values reported for glacier ice, as given here in Table 195. They remark:

Table 195.—Viscosity of Glacier Ice

Adapted from R. M. Deeley and P. H. Parr. 226

		Unit of $\mu_e = 10^{12}$ poises			
Observer		Computer	Computer		
Dr. Main	1888	R. M. Deeley	1912	6.0	
McConnel and Kidd	1888	R. M. Deeley	1912	84.5	
B. Weinberg	1907	B. Weinberg	1907	8.0	
Blumcke and Hess	1907	B. Weinberg	1906	17.4	
Tyndall and others		R. M. Deeley	1908	78.9	
Blumcke and Hess	1910	B. Weinberg	1910	17.5	
Blümcke and Hess	1910	Deeley and Parr	1913	147.70	
Blümcke and Hess	1910	Deeley and Parr	1913	125ª	

From motion of glaciers in the winter.

[&]quot;We have seen that glacier ice consists of crystal granules which not only shear freely along planes at right angles to the optic axis, but also undergo

changes at their bounding surfaces, which enable the mass to suffer continuous distortion under stress. The ability of glacier ice to spread out into piedmonts whose upper surfaces are very nearly level also shows that such shear may take place under very small stresses." This accords with the small value of the shear (λ) that corresponds to the elastic limit. From Table 191, we find that λ does not exceed 60 microradians, nor does the coefficient of rigidity exceed about 3×10^{10} dynes/cm²-radian; hence ice will yield continuously if the shearing stress exceeds 1.8 megadynes/cm² = 1.8 kg*/cm², a very small value.

For other values of the viscosity of glacier ice, as derived from the observed flow of each of several glaciers, see R. M. Deeley.²⁶¹

From the damping of torsional vibrations of a cylinder of ice about its axis, C. D. Hargis 240 obtained the values $\mu_e=3.7$ megamegapoises when the period was 0.286 sec, and 6.21 when it was 0.448 sec. The cylinder was obtained by freezing water in a brass tube.

All the preceding may be summarized thus: (1) None of the available data for the plasticity or for the viscosity of ice is entirely satisfactory. (2) Values of μ_0 derived from the bending of bars are of the order of 10^{10} poises, those from the axial torsion and those from the longitudinal stretching of bars are of the order of 1012 poises. (3) Although McConnel's and McConnel and Kidd's data indicate that μ_e for shear parallel to the optic axis is about 100 times as great as for shear perpendicular to that axis, Hess's data indicate that the difference is slight. (4) The value of μ_0 increases as the rate of shear decreases (Table 193). (5) When the stress is kept constant, μ_e increases with the time the stress has been applied. Whether this involves other phenomena than those pertaining to the variation with the rate of shear cannot be determined from the data now available. (6) The value of μ increases very rapidly as the temperature decreases, a decrease of 10 °C being accompanied by a 5-fold increase in μ for river ice, and a 26-fold increase for glacier ice. This increase in μ causes a marked, but in general a smaller, increase in μ_e . (7) An attempt to fit the data of Hess, of McConnel, and of McConnel and Kidd to Weinberg's equation has been unsuccessful. (8) Owing to the absence of important data, to significant variations in the procedures followed, and to variations in the structure and the purity of the ice used, it is impossible to correlate satisfactorily the data obtained by different observers.

Sustaining power of an ice sheet.—A knowledge of the load that a sheet of ice of given thickness can sustain while resting upon water is of considerable importance, especially in military operations. Ordinary experience teaches, as pointed out by F. A. Forel,²⁶² that this load depends upon the state of the ice. Old ice that has been exposed to the sun and to air not much below 0 °C becomes split by a multitude of vertical cracks into irregular prismatic needles, i.e., it becomes rotten. Such ice has little sustaining

²⁶¹ Deeley, R. M., Geol. Mag. (5), 9, 265-269 (1912).

ses Forel, F. A., Rev. Sci., 51, 379 (1893).

power; no estimate of that power can be given, as it varies greatly with the existing condition of the ice. It is only of new ice still in the process of formation that numerical data can be given with any confidence. It is to such ice that the following figures refer.

An anonymous note ²⁶³ quotes from the "Echo de l'Armée" the following values as having been determined under the authorization of the French military establishment: When 4 cm thick, such ice will bear the weight of one man; when 9 cm thick, infantry marching in open formation; when 12 cm, artillery train of 8-cm guns; when 14 cm, train of 12-cm guns; when 16 cm, siege guns with loaded caissons; when 29 cm thick, it will carry almost any load that would be placed on it.

P. Vedel ²⁶⁴ has stated that the "army rules" were as follows: 2-in. (5-cm) ice will support a man or properly spaced infantry; 4-in. (10-cm) ice, a man on horseback, cavalry, light guns; 6-in. (15-cm) ice, such field-pieces as 80-pounders; 8-in. (20-cm) ice, battery of artillery with carriages and horses, but not over 1000 lb* per sq. ft. on sledges; 10-in. (25-cm) ice, an army, an innumerable multitude; 15-in. (38-cm) ice, railroad tracks and trains. He stated that 24-in (61-cm) ice withstood the impact of a loaded railroad passenger car falling 60 ft (which he estimated at 1500 ft·tons*), but broke under the impact of a locomotive and tender (which he estimated at 3000 ft·tons*). Tables purporting to give the maximum safe load for the ice on a circular lake and for that on a canal are included in the article, but as the value he gives for the Young's modulus in an earlier portion of the paper is about 1000 times too great, it is feared that the data of those tables are untrustworthy. The present compiler has not attempted to check the computations.

65. Deformability of Snow

The deformability of snow and its variation with the depth of the overlying snow have recently been studied by M. Kuroda.²⁶⁵

For obtaining an estimate of the hardness, he used a brass-tipped wooden cone, vertex angle = 90° , dropped from a stated height, and measured (D) the surface diameter of the indentation produced.

For measuring the tensile strength he used telescoping sheet-metal forms of the general shape of the axial section of metal specimens intended for similar tests. One of these forms was pressed into a layer of undisturbed snow carefully taken up on a glass plate, and the force required to pull it apart was measured.

For obtaining the shearing strength he used a flat block sliding snugly in a slot cut in another block; through the center of the compound block and perpendicular to the plane of the sliding one was cut a rectangular hole. By means of a suitable sheet-metal form, prisms of snow that fitted

²⁰⁸ Anon., Idem, 51, 318 (1893).

²⁰⁴ Vedel, P., J. Franklin Inst., 140, 355-370, 437-455 (1895).

²⁰⁰⁵ Kuroda, M., Sci. Papers Inst. Phys. and Chem. Res. (Tokyo), 12, 69-81 (1929).

Table 196.—Hardness of Snow: Variation with Depth in Snow-blanket Adapted from M. Kuroda. 265

The hardness is indicated by the surface diameter (D) of the conical indentation produced by dropping a brass-tipped wooden cone, vertex angle = 90°, from a stated height (h) above the surface of snow under study. By carefully removing the overlying snow, that surface was placed at any desired depth (d) below the undisturbed surface of the natural snowblanket; h=0 indicates that the cone was placed gently upon the surface and sank under its own weight. Values are given for two blankets; those in the first column h=0 refer to one, and the others, to the other. In the original paper the values of h and d for the second blanket appear to have been interchanged, they are the reverse of those here given.

Unit of h , d , and $D=1$ cm										
$h \rightarrow$	0	0	5	10	20					
d	D			_ D						
0	16.8		16.0	19.2						
2	14.0									
10	10.0		4.7	9.1	11.7					
15	4.4									
20	4.3		4.6	8.0	12.0					
30	4.0									
40	4.5	3.4		7.6	12.0					
50	4.0									
60	5.0									
70	3.3	3.0		5.4	8.0					
80	3.6									

Table 197.—Hardness of Snow: Effect of Tamping 265

By carefully removing the overlying snow, the surface under study could be brought to any desired depth (d) below the surface of the natural snow-blanket. The hardness of one portion of the surface so cleared was determined at once; another portion was tamped by a single dropping from a height H of a load of 2700 g* with a rectangular base 18 by 27 cm, and the hardness was then determined. D and h have the same significance as in Table 196.

Two snow-blankets were studied; temperature of snow, about -11 °C.

		h = 20 cm.	Unit of H , d , $D=1$ cm		
H→ d 0 10	L'a	20	Į'a	40 D	60
d	4.5	- D	140	<i>D</i>	
U	4.5		14.0		
10	4.5 8.5		8.0		
12	8.5	4.0			
13			5.5	2.0	1.5
16	6.0	3.0			
19	6.5	3.0			
20	5.2		4.5	2.5	2.0
22	5.2	4.8			
20 22 25 26 30	4.5		4.0	3.3	3.0
26	4.0	3.5			
30	3.0	3.0	3.5	3.5	3.5
45	4.5	4.5			

U = untamped.

the hole could be cut out and placed in it; the force then required to withdraw the sliding block was measured.

He gives curves showing the grain size, the density, D, and the temperature, throughout the thickness of a natural snow-blanket 9 meters thick. The size of the grains varied but little until the ground was approached, where the snow was several months old; the density varied from 0.35 g/cm³ at a depth of 50 cm to 0.65 at 700 cm; from a depth of 200 cm to that of 700 cm the hardness was essentially constant; the temperature was lowest $(-0.9\,^{\circ}\text{C})$ at mid-depth.

Table 198.—Strength and Hardness of Snow 265

T = tensile strength, S = shearing strength, D = hardness as in Table 196, t_8 °C = temperature of the snow. For details, see text.

Unit of $D=1$ cm; of T and $S=1$ g*'cm ²								
Snowa	A	В	C	D				
t.	-9.0	0	-2.0					
T	63	33	93					
S	3	2.5	20	43				
D	20	20	155					

[&]quot;Snow: A= fresh and powdery; B= wet and soft; C= surface crusted; D= surface more crusted than C.

66. ACOUSTIC AND OTHER VIBRATIONAL DATA FOR ICE

(For the elastic constants of ice, see Sections 64 and 67; for density, Section 67.)

Acoustic data are those pertaining to longitudinal vibrations, to those in which the displacement is in the direction of propagation of the train of waves. Like other solids, ice can transmit transverse vibrations also, those in which the displacement is perpendicular to the direction of propagation; and a thin sheet of ice can transmit flexural vibrations. All these types of vibration are considered in this section.

Velocity of Transmission.

As ice is crystalline, the velocity of a given type of vibration might be expected to vary with the direction of propagation through the crystal, and R. Köhler ²⁶⁶ thought that his observations on an ice sheet 30 cm thick indicated such an effect for waves generated by the firing of explosives. But R. W. Boyle and D. O. Sproule ²⁶⁷ concluded, from their observations on longitudinal ultrasonic waves in rods of ice, that any such difference lies within the experimental error (see Table 199). It would seem that their observations are the more readily interpretable.

Flexural waves in an ice sheet 11 to 38 cm thick, resting on water 1.2 to 6 m deep, exhibit marked dispersion; the group velocity is $12.2\sqrt{\tau v}$

²⁰⁰ Köhler, R., Z. Gcophys., 5, 314-316 (1929).

²⁰⁷⁷ Boyle, R. W., and Sproule, D. O., Can. J. Res., 5, 601-618 (1931).

m/sec where τ cm is the thickness of the ice, and the frequency is ν cycles/ sec, v varying from about 13 to 600.268

Table 199.—Velocity of Waves in Ice

(For flexural waves, see text.)

 θ = angle between the direction of advance of the wave and the normal to the planes of freezing (the optic axis); $\nu =$ frequency; $\nu =$ velocity of propagation. The BS^a observations for $\theta = 0$ are given within 0.9 per cent by v = 3.12 (1 - 0.0025t) km/sec, which for t = -26 °C is 3.33; the differences between this and the several values tabulated for -26 °C represent inherent variations in the samples.

Unit of v = 1 km/sec; of v = 1 kilocycle/sec. Temp. = 1 °C

I. Longitudinal vibrations. Rods of ice.

	BS ^a	BS^a $t = -26 ^{\circ}C$		RS^a $t = -4 ^{\circ}C$		ECT ^a			
•	$=0^{\circ}$ = 13		to 12		1.55	y = 1	1.31 to 4.97		
1	\boldsymbol{v}	θ	υ	θ	v	θ	v		
- 9	3.18	0°	3.22	90°	3.23	90°	3.174		
-10	3.21	90°	3.24			$Vb1^b$	3.150		
-30	3.33	90°	3.18¢			$Mean^b$	3.163 ± 0.009		
_ 35	3 43	450	3 11						

11. Longitudinal. Explosions. III. Transverse. Explosions.

Form	v	Ref.ª	Form	v	Ref.a
Sheet	3.40d	ECT	Rod	1.914 ± 0.006	ECT
Sheet	3.41	ECT	Sheet	1.846 ± 0.005	EC
30 cm Sht.	3.23	K	Sheet	1.70	K
Solid	3.15^{d}	ECT	Glacier	1.60	\mathbf{M}
Glacier	3.40	M	Glacier	1.69	\mathbf{M}
Glacier	3.60	M	Glacier	1.67	\mathbf{M}
Glacier	3.57	\mathbf{M}	Glacier	1.69	S
Glacier	3.49	S	Glacier	1.60	S S
Glacier	3.41	S	Glacier	1.82	
Glacier	3.70	S	Névé	1.35	\mathbf{M}
Névé	3.14	\mathbf{M}			

a References and notes:

R. W. Boyle and D. O. Sproule 207; rods cut from river ice. BS

EC M. Ewing and A. P. Crary.208

M. Ewing and A. P. Crary.

M. Ewing, A. P. Crary, and A. M. Thorne, Jr. 200; rods cut from river ice; explosions in the ice sheet; temperature -5 to -15 °C.

R. Köhler 200; lake ice 30 cm thick.

H. Mothes. 270

M. Reich and O. Stierstadt 271; ice formed from distilled water.

E. Sorge 272; inland ice sheet, Greenland. ECT K M

RS

Vbl indicates that the ice was formed by freezing water slowly in a vertical brass tube; under which conditions, the crystals are variously oriented. Mean = mean of all observations on rods of whichever kind.

Direction of propagation was perpendicular to that in preceding case, but in

both it was parallel to the planes of freezing.

^d Computed from the observed velocity in thin rods. Sheet = thin plate, unsupported. Solid = infinite solid.

• Torsional vibrations of thin rod of Vbl ice (see note •), $\nu = 0.81$ kcycle/sec.

²⁰⁰ Ewing, M., and Crary, A. P., Physics, 5, 181-184 (1934) → Crary and Ewing, Phys. Rev. (2), 45, 749 (A) (1934).

Reflectivity.

As compared with steel, or even with granite, ice is a very poor reflector of ultrasonic vibrations ($\nu = 84$ kcycle/sec). See R. W. Boyle and G. B. Taylor.²⁷⁸

67. Pressure-Volume-Temperature Associations for Ice

All data pertaining to the specific volume and the density of ice and of snow, and to their variations with the temperature and the hydrostatic pressure, are assembled in this section. The density has not been directly determined for any type of ice except the usual one (ice-I), and no determination of the density of snow-crystals has come to the compiler's attention. (Deformability of ice and of snow, see Sections 64 and 65. Linear expansion of ice, see Section 68.)

Density of Snow.

The density of freshly fallen snow varies greatly, depending upon the aerodynamic conditions attending its deposition; the density at any point of a snow blanket increases with the age of the blanket, even in the absence of fusion.²⁷⁴ The density in a natural blanket of snow increases nonlinearly with the depth.^{234, p. 25} M. Kuroda ²⁷⁵ found 0.35 g/cm³ at a depth of 50 cm, and 0.65 at 700 cm. Values as low as 0.004 have been recorded for freshly fallen snow.²⁷⁶

The density of the persistent $n\acute{e}v\acute{e}$ in the Pyrenees at altitudes of 2.5 to 3.4 km varies from 0.51 to 0.59 g/cm³ in August to September, and from 0.53 to 0.65 in October.²⁷⁴ Devaux thought that this apparent increase was real. The samples were probably taken from near the surface. E. Sorge ²⁷⁷ has found that the density of the $n\acute{e}v\acute{e}$ on the inland ice-sheet of Greenland is 0.51 g/cm³ at depths of 30 to 118 cm, varying inappreciably with the depth.

Density of Ice-I at 1 Atmosphere.

The density of the ice of glaciers is, as one would expect, lower than that of clear compact ice; values varying from 0.86 to 0.91 g/cm³ have been reported by J. Devaux.²⁷⁸

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200 Ewing, M., Clary, A. P., and Thorne, A. M., Jr., Physics, 5, 165-168 (1934) → Phys. Rev. (2), 45, 749 (A) (1934).

270 Mothes, H., Z. Geophys., 3, 121-134 (1927); 5, 120-144 (1929).

271 Reich, M., and Stierstadt, O., Physik. Z., 32, 124-130 (1931).

272 Sorge, E., Z. Geophys., 6, 22-31 (1930).

273 Boyle, R. W., and Taylor, G. B., Trans. Roy. Soc. Canada, III (3), 20, 245-257 (1926).

274 Devaux, J., Compt. rend., 185, 1147-1149 (1927).

275 Kuroda, M., Sci. Papers Inst. Phys. & Chem. Res. (Tokyo), 12, 69-81 (1929).

276 Keränen, J., Annales Acad. Sci. Fennicae = Suomalaisen Tiedeak. Toimit. (A), 13, No. 8 (1920).

277 Sorge, E., Z. Geophys., 6, 22-31 (1930).

278 Devaux, J., Compt. rend., 185, 1602-1604 (1927).
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The better of the recorded values for the density of ice-I at 0 °C and 1 atm vary from 0.918 to 0.916 g/ml. E. L. Nichols ²⁷⁹ has reported values indicating that the density is actually subject to such variations. His values indicate that the density of freshly formed natural ice is 0.91795 g/ml; of natural ice 1 year old, 0.91632; and of artificial ice-mantles frozen by means of solid CO₂ and ether, 0.91603. But H. T. Barnes, ²⁸⁰ using ice from the St. Lawrence River, failed to find such great variation. He found for new ice 0.91662, for ice 1 year old 0.91648, and for ice 2 years old 0.91637, the mean of all being 0.91649 g/ml. Although these values indicate a slight progressive decrease in the density as the ice ages, the change in 2 years is only about 1/6 of that reported by Nichols for 1 year. J. H. Vincent ²⁸¹ also believes that he has shown "that the same specimen of water may assume different densities on freezing." The values he finds vary from 0.9155 to 0.9163.

At least a portion of the differences in the densities recorded is probably due to the fact that the water from which the ice was formed contained other substances in solution. J. Y. Buchanan ²⁸² has shown that, when ice forms in an aqueous solution, some of the solution is entrapped between the crystals, and remains incompletely frozen so long as the temperature is above the cryohydric point. This makes the apparent density too great. The effect may be appreciable even when the solution is extremely dilute, as dilute as good distilled water. (See Table 203.) The pressure caused by the expansion attending the freezing of a portion of a volume of water entirely surrounded by ice will tend to keep the remainder of the volume in the liquid state, if the temperature is not far from zero. This also increases the apparent density.

The subject of the variability in the apparent density of ice should be given careful study, using ice of various ages, from various sources, and frozen under various controlled and recorded conditions. Special attention should be given to the possible effect of supercooling, of the presence of air, and of the size of the grains and its progressive increase, as reported by R. Emden,²⁸³ not entirely forgetting the so-called "dense ice" reported by Cox (1904) and by Shaw (1924) (see Section 57). The effects of stresses, arising either from a difference in the expansions of ice and its container or from the expansion that occurs on freezing, should be considered more carefully than they have been in the past. It might be worth while trying to use monocrystals, either grown directly or obtained by disintegrating a carefully frozen sheet of ice by means of radiation. And one should not forget the suggestion ²⁸⁴ that variations in the density may

²⁷⁰ Nichols, E. L., Phys. Rev., 8, 21-37 (1899).

²⁸⁰ Barnes, H. T., *Idem*, 13, 55-59 (1901) \rightarrow *Physik. Z.*, 3, 81-82 (1901) = Barnes, H. T., and Cooke, H. L., *Trans. Roy. Soc. Canada, III* (2), 8, 143-155 (1902).

²⁸¹ Vincent, J. H., *Phil. Trans.* (A), 198, 463-481 (1902) \rightarrow *Proc. Roy. Soc. (London)*, 69, 422-

<sup>424 (1902).

282</sup> Buchanan, T. V. Proc. Roy. Soc. Edinburgh, 14, 129-149 (1887); Nature, 35, 608-611 (1887);

²⁸² Buchanan, J. Y., Proc. Roy. Soc. Edinburgh, 14, 129-149 (1887); Nature, 35, 608-611 (1887); 36, 9-12 (1887); Proc. Roy. Inst. Grt. Britain, 19, 243-276 (1908).

²⁸⁸ Emden, R., Neue Denkschr. d. allgem. schweiz. Ges. Naturwiss., 33, 43 pp. (1892).

²⁸⁴ Emeléus, H. J., et al., J. Chem. Soc. (London), 1934, 1207-1219 (1934).

arise from actual changes in the composition of the water, such as a concentration of deuterium oxide during the process of freezing; even though the possibility of so concentrating deuterium oxide seems to have been disproved by the work of V. K. LaMer, W. C. Eichelberger, and H. C. Urey,²⁸⁵ and of G. Bruni.²⁸⁶ In the last two of these papers, Bruni disproves the contrary conclusion drawn in an earlier paper by G. Bruni and M. Strada.²⁸⁷

The more reliable determinations of the density of ice are listed in Table 200, where Kopp's distinctly abnormal value is also given. Those for natural and for artificial ice have been listed separately, simply because it has been suggested that they may differ. No such difference is at all obvious from the table.

The values may, however, be assorted into three distinct groups. One group includes the values obtained by C. Brunner ²⁸⁸ for river ice, by E. L. Nichols ²⁷⁹ for new ice, by L. Dufour ²⁸⁹ for ice from boiled-out water frozen in air at a very low pressure, and by A. Leduc ²⁹⁰ for ice from which the air had been removed with great care. The average of these values is 0.9178, and the average deviation from this is 0.0002.

At the other extreme is the group of values obtained from direct measurements of the difference in the specific volumes of ice and of water. This contains the unsatisfactory determination of H. Kopp,²⁹¹ the essentially identical values (0.9157) obtained by Plücker and Geissler ²⁹² and by H. Endo,²⁹³ and the slightly higher ones (0.9160) found by J. H. Vincent ²⁸¹ and by Nichols ²⁷⁹ for artificial ice. The mean, omitting Kopp's value, is 0.9158.

The others fall into the third group, averaging 0.9165, and ranging from 0.9163 to 0.9166, the mean variation being about 0.0002.

The reason for such variations and for such a grouping of the values remains to be determined.

The values have usually been reported in terms of the density of water at 0 °C, and consequently differ slightly from the values here given. Still other changes occasionally seemed justified. They are described in the following remarks concerning the several determinations.

J. Plücker and Geissler ²⁹² concluded that whenever the freezing occurs in the same way the density of the ice is always the same. They used a unique type of thermometer having in the bulb a distinct, but communicating compartment for the water under study. The value tabulated has

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<sup>285</sup> LaMer, V. K., Eichelberger, W. C., and Urey, H. C., J. Am. Chem. Soc., 56, 248-249 (1934).

<sup>286</sup> Bruni, G., Att. d. R. Acc. Naz. Lincei (6), 20, 73-75 (1934)—J. Am. Chem. Soc., 56, 2013-2014 (1934).
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²⁸⁷ Bruni, G., and Strada, M., Ibid, 19, 453-458 (1934).

²⁸⁸ Brunner, C., Ann. d. Phys. (Pogg.), 64, 113-124 (1845).

²⁸⁰ Dufour, L., Compt. rend., 50, 1039-1040 (1860); 54, 1079-1082 (1862).

²⁹⁰ Leduc, A., Idem, 142, 149-151 (1906).

²⁹¹ Kopp, H., Ann. d. Chem. u. Pharm. (Liebig), 93, 129-232 (1855) \rightarrow Ann. de chim. ct Phys. (3), 47, 291-296 (1856).

²⁰² Plücker, J., and Geissler, Ann. d. Phys. (Pogg.), 86, 238-279 (265-279), (1852).

²⁹³ Endo, H., Sci. Rep. Tôhoku Imp. Univ. (Sendai) (1), 13, 193-218 (1924-25).

been derived from their determinations of the difference between the specific volumes of ice and water. With one exception, omitted in deriving the mean, their observations are exceedingly concordant.

- H. Kopp ²⁹¹ measured the expansion that occurs when water freezes, using turpentine as the dilatometric liquid. Although his dilatometer was closed with a cork, he obtained two closely agreeing values giving for the density 0.908. Like other determinations by this method, the value is lower than that found by any other method, but this particular value is undoubtedly too low. (In the French abstract the values given for the contraction are wrong.)
- L. Dufour ²⁸⁹ used a flotation method, and ice from boiled-out water frozen in a vacuum (air-pressure not over 0.5 mm-Hg). In the earlier work he used a mixture of alcohol and water, and found that a density of 0.9175 g/cm³ just below 0 °C was required to support completely submerged ice. This mixture was found to dissolve the ice slightly. In the later work, he used a mixture of chloroform and petroleum, worked between -0.5 °C and -8 °C, and accepted 0.000158 (°C)-1 as the coefficient of cubical expansion of ice. Sixteen determinations of the density of ice at 0 °C lay in the range 0.9168 to 0.9193 and averaged 0.9178.

The observations of R. Bunsen 294 vary linearly with the temperature (τ) at which the water was frozen. They may be represented by the formula $d=0.91663-0.000047\tau$; the lower the temperature of freezing, the greater the density. In how far this arises from the difference in the expansion of ice and of glass remains to be determined. The value corresponding to $\tau=0$ is the one here tabulated.

- J. v. Zakrzewski ²⁹⁵ used Bunsen's method, but froze the water at a constant temperature not far from 0 °C. The mean of three very closely concordant determinations of the density at -0.701 °C, the temperature of freezing, gave $d_{-0.701} = 0.916710$ which reduces to $d_0 = 0.91661$ g/ml if $10^6\beta = 153$ (see Table 202). A single measurement at -4.720 °C gave $d_{-4.720} = 0.916995$ or $d_0 = 0.91633$; if the temperature recorded as -4.720 was actually -2.720, then $d_0 = 0.91661$, agreeing closely with the mean of the three at -0.701 °C. The -4.7 °C value has not been entered in the table.
- E. I. Nichols 279 determined the density by weighing ice in air and in petroleum. The highest value for the specific gravity with reference to water at 0 °C was 0.91808 for a recently formed, natural icicle; the lowest was 0.91615 for an ice mantle frozen by means of ether and solid carbon dioxide, temperature about -70 °C. Other measurements of a tentative kind are described.
- J. H. Vincent.²⁸¹ using a novel device, weighed in mercury the buoyancy of water and of the ice formed from it. The quantity directly determined was the difference in the specific volumes. The buoyancy of the ice was weighed at several temperatures in the range -0.4 °C to -10 °C, and the

²⁰⁴ Bunsen, R., Ann. d. Phys. (Pogg.), 141, 1-31 (1870).

²⁰⁰ v. Zakrzewski, J., Ann. d. Phys. (Wied.), 47, 155-162 (1892).

value at 0 °C was obtained by linear extrapolation. His values for the density at 0 °C vary from 0.915460 to 0.916335; he took as the weighted mean 0.9160. Two freezings of the same specimen of water gave densities differing by 57 parts in 100 000. He regarded this difference as real.

A. Leduc 290 took great pains to remove all air from the water from which the ice was formed. For determining the density, he froze the water in a specific-gravity flask of the Regnault type. The freezing proceeded gradually from the bottom up into the capillary; a mixture of ice and salt was used, the temperature being between -5° and -10° C.

Table 200.—Density of Ice-I at Atmospheric Pressure

(Comments on the several determinations may be found in the text.)

Dewar's (J. 1902) corrected (see text) value for -188.7 °C is $d_{-188.7} = 0.936$; this with the coefficient (Table 202) $10^6\beta = 153$ gives 0.948 for the density at absolute zero (-273.1 °C), which probably exceeds the true value. From $d_0 = 0.9166$ and $10^6\beta = 153$ one derives $d_{-188.7} = 0.943$.

The value accepted by J. R. Clarke ³⁰⁰ for the density at 0 °C is $d_0 = 0.9168 \pm 0.0005$ g/ml (ICT).

 d_t = density at $t \circ \hat{C}$; $\tau \circ C$ is the temperature at which the ice was frozen, value for $\tau = 0 \circ C$ is obtained by extrapolation.

of $d = 1$ g/ml = 0.999973 g/c	m ³	
	Artificial Ice	
ge Observer	d_0	Notes
Plücker and Geissl	er	
(1852)	0.9156_{7}	Ch. in vol.
w ^a Kopp (1855)	0.908	Ch. in vol.
r. Dufour (1860)	0.9175	Flotation
ew Dufour (1862)	0.9178	Flotation
r. Bunsen (1870)	0.91663	$\tau = 0$
	0.91661	$\tau = -0.701$
Nichols (1899)	0.91603	$\tau = -70$
Vincent (1902)	0.9160	Ch. in vol.
Leduc (1906)	0.9176	$\tau = -5$ to -10
Endo (1925)	0.9157_{1}	Ch. in vol.
	Observer Plücker and Geissl (1852) Wa Kopp (1855) Or. Dufour (1860) Ow Dufour (1862) Or. Bunsen (1870) Or. V.Zakrzewski (1892) Nichols (1899) Vincent (1902) Leduc (1906)	Observer Artificial Ice Plücker and Geissler (1852) 0.9156, Wa Kopp (1855) 0.908 Ort. Dufour (1860) 0.9175 Ow Dufour (1862) 0.9178 Ort. Bunsen (1870) 0.91663 Ort. V.Zakrzewski (1892) 0.91661 Nichols (1899) 0.91603 Vincent (1902) 0.9160 Leduc (1906) 0.9176

[&]quot;Mean of three values: Icicles 0.91804, 0.91789, and new pond ice 0.91792 g/ml.

- H. Endo ²⁹³ determined the change in specific volume from observations of the buoyancy, in lamp oil (kerosene?) at various temperatures, of a silica vessel containing the specimen. The ice was frozen by means of a mixture of ether and solid carbon dioxide.
- J. Dewar ²⁹⁶ weighed in air and in liquid air "pieces of clear ice cut from large blocks." The density of the liquid air was determined in terms of that of liquid oxygen boiling under a pressure of 1 atm, assumed to be 1.137. Thus he obtained for ice at -188.7 °C the value 0.930₀, individual

²⁸⁰ Dewar, J., Proc. Roy. Soc. (London), 70, 237-246 (1902) → Chem. News, 25, 277-279, 289-290 (1902). Same data in Proc. Roy. Inst. Grt. Brit., 17, 418-426 (1903) → Chem. News, 91, 216-219 (1905).

values ranging from 0.926₅ to 0.933₂. Reducing this value to the basis of 1.144₇ ²⁹⁷ for the density of oxygen under the stated conditions, we find 0.936₀. It is interesting to note that this is only 0.75 per cent less than

Table 201.—Densities and Specific Volumes of the Ices at their Melting-points.

(For change in volume on transition of ice to ice, see Table 271.)

The following data have been obtained by combining those (Bridgman's) in Tables 95 and 271, interpolating or extrapolating the water data where necessary.

 $d = \text{density}, v^* = \text{specific volume}, mp \, ^{\circ}\text{C} = \text{melting point corresponding to } P.$

Unit of P = 1 atm = 1.01325 megadyne/cm²; of d = 1 g/ml; of $v^* = 1$ ml/g Type of Ice 21* dw vw* dι mp 104(vi* - vw*) I 0.0 0.9168 1 1.0908 0.9921 1.0008 +900-5.0590 0.9740 0.9297 1.0756 1.0267 +1016-10.01090 0.9397 1.0642 1.0504 0.9520 +1122-15.01540 0.9444 1.0589 1.0671 0.9371 +12180.9481 -20.01910 1.0547 1.0830 0.9234 +1313 -22.0° 2045 0.9483 1.0545 1.0878 0.9193 +1352Ш -17.03420 1.1595 0.8624 1.1293 0.8855 -231 -17.0° 3420 1.1609 0.8614 1.1293 0.8855 -241-18.52820 0.8686 0.8987 -3011.1513 1.1127 0.8714 -20.02430 1.1476 1.1101 0.9085 -371-22.0°2045 1.1459 0.8727 1.0878 0.9193-466V 0.16ª 6175 0.7901 -5271.2657 1.1865 0.84280.7903 0.0 6160 1.2653 1.1862 0.8430-527-5.05270 1.2596 0.7939 1.1707 0.8542 -6034360 1.2488 -10.00.80081.1511 0.8687 -679 -15.03680 0.8051 1.1357 0.8805 -7541,2421 0.8067 0.8855 -788 -17.0° 3420 1.2396 1.1293 1.2338 -20.03040 0.8105 1.1194 0.8933 -82811 990 0.7344 1.2604 0.7934 -590VI +40.01.3616 0.8055 +30.010 250 1.3528 0.7392 1.2415 -663 $+20.0^{\circ}$ 8710 1.3492 0.7412 1.2250 0.8163 -751 +15.01.3464 0.74271.2158 0.8225 -- *7*98 8040 +10.07390 1.3430 0.7446 1.2063 0.8290-844 0.7459 1 1986 -884+5.06880 1.3407 0.8343 0.7512 1.1865 $+0.16^{\circ}$ 6175 1.3312 0.8428 -916 0.7514 -9161.1862 0.8430 0.0 6160 1.3308 0.7555 -9381.1774 0.8493 -- 5.0 5620 1.3236 -10.0-960 1.3158 0.7600 1.1682 0.8560 5110 1.159 0.863 -980-15.04640 1.308 0.765 VII 20.0 48 400 1.67 0.60

⁶ At 25.0 °C the melting pressure is 9630 bars (=9504 atm) and $v_i^* - v_w^* = -0.0714$ cm⁸/g, Bridgman's value being -0.0707.

"The data for water needed for the computation of the densities are not available; values of $(v_i^* - v_w^*)$ are given in Table 271. This value for v_i is from P. W. Bridgman.

[&]quot;Triple point, water and two ices.

²⁹⁷ ICT, Int. Crit. Tables, 3, 20 (1928).

that computed on the assumption that the mean coefficient of expansion between 0 °C and -190 °C is equal to the coefficient (153 × 10 °) found near 0 °C.

No comments need be made regarding the other determinations; references for them have already been given. For discussions of the experimental values, see E. L. Nichols,²⁷⁹ H. T. Barnes and H. L. Cooke,²⁹⁸ and W. A. Roth.²⁹⁹

Densities of the Ices not at their Melting-points.

In addition to values that may be derived from the compressibilities, the coefficients of thermal expansion, and the specific volumes, the following values have been reported.

- Ice-II. From x-ray data for ice-II at -155 °C and atmospheric pressure the density 1.21 g/cm³ has been derived.³⁰¹
- Ice-III. From x-ray data for ice-III at -155 °C and atmospheric pressure McFarlan has concluded that the density is $1.103 \text{ g/cm}^3.302$
- Ice-VI. L. H. Adams ³⁰³ has reported the following values for the specific volumes of ice-VI (v_i^*) and of water (v_w^*) , each at 25.0 °C and under the indicated pressure:

P_b	<i>7</i> 000	8000	9000	10,000	11,000	12,000	bars
P	6890	7895	8882	9869	10,856	11,843	atm
10⁴vw*	8402	8278	8166	8059	7964	7876	cm ³ /g
104716*	7509	7463	7417	7371	7325	7279	cm ⁸ /g
d_{i}	1.3317	1.3399	1.3482	1.3567	1.3652	1.3738	g/cm³
			(See also ?	Γable 201).			

Thermal Expansion of Ice (Cubical). (For linear expansion, see next Section.)

In the reports of the *Vega* expedition, O. Pettersson ³⁰⁵ gives data indicating that the thermal coefficient of cubical expansion of ice is about 170×10^{-6} per 1 °C if the temperature is not above -3 °C, and decreases as the temperature rises above that point, becoming negative (*i.e.*, there is a contraction) as the melting point is closely approached. The purer the water from which the ice is frozen, the smaller is the variation in the coefficient, and the nearer to the melting point does the contraction first appear.

This strange behavior, which may account in part for the conclusion of O. Fort 306 that the early observations of A. Petzholdt 307 indicated that ice

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298 Barnes, H. T., and Cooke, H. L., Trans. Roy. Soc. Canada, III (2), 8, 143-155 (1902).
290 Roth, W. A., Z. physik. Chem., 63, 441-446 (1908).
200 Clarke, J. R., Int. Crit. Tables, 3, 43 (1928).
201 McFarlan, R. L., J. Chem'l Phys., 4, 60-64 (1936) → Phys. Rev. (2), 49, 199 (A) (1936).
202 McFarlan, R. L., J. Chem'l Phys., 4, 253-259 (1936) → Phys. Rev. (2), 49, 644 (A) (1936).
203 Adams, L. H., J. Am. Chem. Soc., 53, 3769-3813 (1931).
204 Bridgman, P. W., J. Chem'l Phys., 5, 964-966 (1937).
205 Pettersson, O., Beibl. Ann. d. Physik, 7, 834-841 (1883).
206 Fort, O., Ann. d. Physik (Pogg.), 66, 300-302 (1845).
207 Petzholdt, A., "Beiträge zur Geognosie von Tyrol," 1843.
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expands as the temperature falls, is thus accounted for by J.Y. Buchanan ²⁸²: The ice formed from a dilute solution contains no salt, but some of the solution is entrapped between the crystals. Ice separates from that solution until the concentration of the solution has risen to such a value that the existing temperature is that at which there is equilibrium between the solution and ice. Thus the volume of liquid enclosed by the ice varies with the temperature, but is never zero so long as the temperature is above the cryohydric point. The actual change in volume as the temperature rises is the resultant of two effects: (a) the expansion of the ice; (b) the contraction attending the melting that is required to adjust the concentration of the entrapped solution. Above some temperature, the latter overbalances the former. This explanation assumes that the overall volume of a block of ice follows the changes in the volume of the entrapped material.

Table 202.—Isopiestic Coefficient of Cubical Expansion of Ice

For ice-VI, P. W. Bridgman ³¹² has stated that within the range 0 to $20 \,^{\circ}\text{C} \, (\partial v^*/\partial t)_p = 120 \times 10^{-6} \, (\text{cm}^3/\text{g})$ per $1 \,^{\circ}\text{C}$.

Values for ice-I are tabulated below; $\beta = \frac{1}{v} \left(\frac{\partial v}{\partial t} \right)_p$; with the exception of Pettersson's observations (see text), β seems to be essentially independent of t; the pressure is nominally 1 atm.

Unit o	of $\beta = 10^{-6}$ per 1 °C.	Temp. = t °C	
Observer	Year	β	Range of t
Brunner 810	1845	122	-0.8 to -19.5
Plücker and Geissler 202	1852	155^{a}	0 to -24
Pettersson 805	1883	170	t below -3
v. Zakrzewski 206	1892	77	-0.7 to -4.7
Vincent 281	1902	152	-0.4 to -10
3 × linear coefficient °		155	0 to -20
Value of choice		153	

^a In taking the mean, one value (170) has been omitted. ^b See p. 468

He showed, by calculation, that the contraction observed by Pettersson in the case of ice formed from ordinary distilled water is the same as that which arises in the manner just indicated when the ice is formed from a solution of sodium chloride that is so dilute as to contain only 7 grams of Cl per 10⁶ grams of water (Table 203). He also calculated, for various concentrations, the temperature at which the apparent volume of the ice is a maximum, and the amount of liquid then contained in it (Table 203). If the solution contains 1 part of Cl to 10⁶ parts of water, the maximum apparent volume occurs at nearly a quarter of a degree below 0 °C; this solution is "in the category of distilled waters."

Apparently, no other investigator of the density or of the thermal dilatation of ice has either recorded an apparent contraction of ice as its rising

[°] See Table 204.

Table 203.—Specific Volume of Ice from Dilute Solutions Adapted from J. Y. Buchanan. 313

The solute is NaCl. C = concentration of the solution at $0 \, {}^{\circ}C$; $W_0 =$ volume at $0 \, {}^{\circ}C$ of the solution that goes to the formation of ice of volume V at the indicated temperature $(t \text{ or } t_m)$; $w_0 =$ volume at $0 \, {}^{\circ}C$ of the unfrozen brine contained in V. Let β and β_1 denote the coefficients of cubical expansion of ice and of water, respectively; $\rho_0 =$ ratio of the density of ice at $0 \, {}^{\circ}C$ to that of water at the same temperature; and $k \equiv w_0/W_0$; k is independent of W_0 , but depends upon C and t in such a way that $\lambda \equiv kt/C$ varies but slightly. Hence a close approximation to k for any pair of intermediate values of t and C can be obtained from the values of λ tabulated below. $\rho_0 V = W_0 \left[1 - k(1 - \rho_0) + \{\beta + k(\rho_0 \beta_1 - \beta)\}t\right]$; the relative contraction due to incomplete freezing is $(W_0 - \rho_0 V)/W_0 \equiv \Delta$. Taking $\rho_0 = 0.9169$, $10^6\beta = + 153$, $10^6\beta_1 = -194$, we have

$$0.9169V = W_0 \left[1 - 0.0831k + (10^{-6})(153 - 331k)t \right]$$

Replacing k by its equivalent $\lambda C/t$, regarding λ as constant, and solving the equation $(\partial V/\partial t)_{\lambda 0} = 0$, one obtains $t_m = -23.3\sqrt{-\lambda C}$ for the temperature at which V is a maximum.

 $(V/W_0)_P$ = value of V/W_0 observed by Pettersson for ice formed from ordinary distilled water.

(In deriving the following values, Buchanan used for ρ_0 , β , and β_1 , values that differ slightly from those just given; in particular, he took $10^6\beta = 160$, which gives $t_m = -22.75\sqrt{-\lambda C}$.)

Unit of $C=1$	g-(1	per			water = 0.000028		per	kg	of	water;
			of \ - 1°	C/	(1 g-Cl per g wa	itei)				

		C 7		
\overline{t}	100k	V/H',	$(V/W_0)P$	- λ ^a
-0.07	1.000	1.08979	1.08980	100
-0.10	0.700	1.09006	1.09007	100
-0.15	0.467	1 09028	1.09038	100
-0.20	0.350	1.09037	1.09048	100
-0.40	0.175	1.09054	1.09057	100
C	tm	100k	1000∆	— λ ^a
10000	-20.5	5.73	7.51	118
5000	-16.6	3.37	5.46	112
2500	-10.75	2.46	3.60	106
1250	-7.8	1.60_{6}	2.48	100.2
1000	−7.0	1.420	2.22	99.4
500	-4.9	1.000	1.56	98.0
250	-3.5	0.695	1.11	97.3
125	-2.55	0.469_{6}	0.77	95.8
100	-2.3	0.4183	0.70	96.2
10	-0.725	0.1363	0.22	98.8
1	-0.2275	0.0438	0.07	99.7
0.1	-0.0725	0.01377	0.02	99.8
0.01	-0.02275	0.004306	0.01	98.0

These values of λ have been derived from the tabulated values of C, k or t_m , and k, which have been taken from Buchanan's paper.

temperature approaches 0 °C, or considered the effect discussed by Buchanan. That effect (the inclusion of unfrozen liquid) has however been considered, and seemingly observed, by A. W. Smith ⁸⁰⁸ and by H. C. Dickinson and N. S. Osborne ³⁰⁹ in their determinations of the specific heat of ice.

Other determinations of the cubical expansion of ice are those of C. Brunner 310 using natural ice and hydrostatic weighings; of J. Plücker and Geissler 292 using an ingenious double-bulb thermometer; of J. v. Zakrzewski, 295 who, using Bunsen's method for determining the density, obtained for the coefficient a value that is only half as great as that found by others; and of J. H. Vincent 281 using a novel hydrostatic method. (Table 202.)

Sir James Dewar ⁸¹¹ has inferred that the coefficient decreases as the temperature is greatly reduced, from the fact that ice at 0 °C cracks in all directions when dropped into liquid air (-188.7 °C), but ice that has been slowly cooled to -188.7 °C does not crack when it is dropped into liquid hydrogen (-252.7 °C). This conclusion accords with the recent determination of the linear expansion by M. Jakob and S. Erk (1928), see Table 204.

Compressibility of Ice.

Ice-I.—In his compilation, L. H. Adams³¹⁴ gives the value obtained by T. W. Richards and C. L. Speyers ³¹⁵ for the isothermal compressibility of ice-I, namely $\gamma \equiv -\frac{1}{v_0} \left(\frac{\partial v}{\partial \rho}\right)_t = 12 \times 10^{-6}$ per bar at -7 °C and 300 bars. Those authors stated that the value of γ between 300 and 500 bars is probably not over 3 per cent less than it is between 100 and 300 bars. (1 bar = 1 megadyne/cm²; they called it a megabar, thus departing from international custom.)

In contrast to this, P. W. Bridgman 316 computed from his observations that at 0 °C and 1 atm $\gamma=37\times10^{-6}$ per bar, a value 3 times as great as the former. Richards and Speyers 315 state that Bridgman has admitted that his calculated value is untrustworthy, and, on their suggestion that the difference may in part be due to γ having a large temperature coefficient, he recalculated γ obtaining the following values for $10^6\gamma$: 0 °C, 33; -5 °C, 23; -7 °C, 21; -10° C, 19; -15 °C, 18. These are still much greater than theirs. They state that Bridgman agreed with them in thinking that the

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    Smith, A. W., Phys. Rev. (2), 17, 193-232 (1903).
    Dickinson, H. C., and Osborne, N. S., Bull. Bur. of Stand., 12, 49-81 (S248) (1915).
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³¹⁰ Brunner, C., Ann. d. Physik (Pogg.), 64, 113-124 (1845).

³¹¹ Dewar, Sir James, Proc. Roy. Inst. Grt. Britain, 17, 418-426 (1903) = Chem. News, 91, 216-219 (1905).

⁸¹² Bridgman, P. W., Proc. Am. Acad. Arts. Sci., 48, 307-362 (1912).

⁸¹⁸ Buchanan, J. Y., Proc. Roy. Inst. Grt. Brit., 19, 243-276 (251, 257) (1908).

⁸¹⁴ Adams, L. H., Int. Crit. Tables, 3, 49-51 (50) (1928).

⁸¹⁵ Richards, T. W., and Speyers, C. L., J. Am. Chem. Soc., 36, 491-494 (1914).

²¹⁰ Bridgman, P. W., Proc. Am. Acad. Arts Sci., 47, 439-558 (1912).

discrepancy may be due to a softening of the ice just before melting. (Is this a reference to the phenomenon discussed by Buchanan, p. 469?)

Ice-VI.—P. W. Bridgman $^{312, p. 362}$ found for ice-VI the value: $-(dv^*/dp)_t = 4.6 \times 10^{-6} \text{ cm}^3/\text{g}$ per bar, essentially constant throughout the ranges 0 to 20 °C and 6000 to 10,000 kg*/cm². (In the International Critical Tables,† this value is incorrectly recorded as what is here denoted by γ .) Bridgman now thinks that that value is probably too high, but he has not yet obtained consistent values for the compressibility at these high pressures for either ice-VI or water.817

Ice-VII.—Mean compressibility of ice-VII between 45,000 and 50,000 kg*/cm² is about 3/4 of that between 20,000 and 25,000 kg*/cm²; on increasing the pressure from 20,000 to 45,000 kg*/cm² the specific volume of ice-VII decreases by 0.039 cm³/g; at 50,000 kg*/cm² and room temperature the specific volume is about 0.60 cm³/g.³¹⁷

68. COEFFICIENT OF LINEAR EXPANSION OF ICE

As ice is a crystalline substance, it is to be expected that its thermal coefficient of linear expansion $(\alpha \equiv (\partial l/l_0 \partial t)_p)$ where l = length, t = temperature, p = pressure) will vary with the angle between l and the optic axis. No data on the expansion of single crystals of ice have been found, but curves representing such data for zinc, which also crystallizes in the hexagonal system, have been obtained by Grüneisen and Goens, and reproduced by M. Jakob and S. Erk.³¹⁸ They indicate that the expansion of zinc in the direction of the hexagonal axis is much greater than it is transverse to that axis, and that in the latter direction the expansion is negative when the temperature is very low.

The only satisfactory series of determinations of α for ice over a considerable range in temperature is that of Jakob and Erk. They used rods of ice frozen slowly in paper tubes, the freezing proceeding radially from outside in. Polariscopic examination gave no indication of any regular orientation of the axes of the constituent crystals, but the variation of α with t was strikingly similar to that for zinc in which t is perpendicular to the optic axis; whence, the authors concluded that the axes of the constituent crystals had a pronounced radial component.

The only other determinations for specimens in which the crystals were thought to have been oriented fairly uniformly are those of C. A. v. Schumacher, of Pohrt, and of A. Moritz, summed up by W. Struve, and quoted by H. Moseley, and extending the work of W. Struve, they indicated that α is essentially independent of the direction of the optic axis.

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† Vol. 3, p. 50.

Bit Bridgman, P. W., J. Chem'l Phys., 5, 964-966 (1937).

Bis Jakob, M., and Erk, S., Wiss. Abh. Physik-Techn. Reichsanst., 12, 301-316 (1928-29) = Z. gesamt. Kälte-Ind., 35, 125-130 (1928).

Bis V. Schumacher, C. A., Mém. Acad. St. Pétersbourg, Math-Phys. (6), 4, 307-357 (1847).

Bis Pohrt, Published with those of Moritz (1847).
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⁸⁸¹ Moritz, A., *Idem*, 4, 358-384 (1847). 882 Struve, W., *Idem*, 4, 294-306 (1847).

Other frequently quoted determinations are those of E. L. Nichols ³²⁵: of T. Andrews, 326 distributed from 0 to -34.5 °C and represented by $10^{6}\alpha = 88.79 + 3.800t + 0.06654t^{2}$; and the very erratic ones of W. H.

Table 204.—Thermal Coefficient of Linear Expansion of Ice (For coefficient of cubical expansion, see Table 202.)

The only satisfactory determinations are those of Jakob and Erk (1928); the others here given are frequently quoted.

Values in parentheses have been computed by means of the appropriate equation given in the text. The specimen used by Andrews was frozen in a cylinder having a height equal to its diameter (2 ft); A and R indicate that the direction of the observed expansion was axial and radial, respectively. θ = assumed angle between the optic axis and the length (1) of the specimen; Vbl indicates that θ is variable, and that the axes are irregularly distributed.

$$\alpha = \frac{1}{l_0} \left(\frac{\partial l}{\partial t} \right)_p$$
; $\alpha_m = \frac{l_2 - l_1}{l_0(t_2 - t_1)}$; $l_0 = \text{value of } l \text{ at } 0 \text{ °C.}$

Unit of α and of $\alpha_m = 10^{-6}$ per 1 °C. Temp. = t °C

Jakob and Erk (1928).318 t t ŧ α t α 33.9 -1500 52.7 -5045.6 -10016.8

Observer	θ	t_2	t_1	α_m		- Andrew	75 326	
v. Schumacher 319	90°	-12°	-27.5°	51.4	θ	t_2	t_1	am
Pohrt 820	90	-11	-26.8	51.1	Vbl R	0	-8.9	73.6
Moritz 321	0	-1.7	-28.5	51.8	Vbl R	-8.9	-17.8	50.3
Struve 324	Vbl	-1.2	-27.5	53.0	Vbl R	-17.8	-29.5	36.9
Nichols 325	Vb1	-8	-12	54.0	Vbl R	-29.5	-34.5	35.5
Sawver 827	Vbl	0	-18	(53.0)	Vbl R	-1	-28	(51.7)
Sawyer 527	Vb1	-1	-28	(42.8)	Vbl R	0	-18	(61.8)
•					Vbl A	0	-17.8	37.0

Sawyer,³²⁷ quoted by H. T. Barnes, J. W. Hayward, and N. M. McLeod,³²⁸ covering the range -8 to -20 °C, and approximately represented by $10^6\alpha = 69.6 + 1.85t$. Neither of the last two sets accords well with the values obtained by others; each is represented by a graph in the compilation by J. R. Clarke. 329

From x-ray studies at 0 °C and -66 °C, H. D. Megaw 330 has con-

⁸²⁸ Moseley, H., Phil. Mag. (4), 39, 1-8 (1870).

³²⁴ Struve, W., Ann. d. Physik (Pogg.), 66, 298-300 (1845).

⁸²⁵ Nichols, E. L., Phys. Rev., 8, 184-186 (1899).

⁸²⁰ Andrews, T., Proc. Roy. Soc. (London), 40, 544-549 (1886).

⁸²⁷ Sawyer, W. H., Proc. Maine Soc. Civ. Eng., 1, 27 (1911).

Barnes, H. T., Hayward, J. W., and McLeod, N. M., Trans. Roy. Soc. Canada III (3), 8, 29-49 (1914).

⁸²⁰ Clarke, J. R., Int. Crit. Tables, 3, 43-45 (43) (1928).

²⁰⁰ Megaw, H. D., Nature, 134, 900-901 (L) (1934).

cluded that the base (a) and the height (c) of the unit cell of ice have the following values: a = 4.5135A and c = 7.3521A at 0 °C, and a = 4.5085A and c = 7.338A at -66° C. These give for the mean coefficient of linear expansion (α_m) between these temperatures the values $10^6\alpha_m = 17$, for a and 29 for c, which are much smaller than would be inferred from any of the measurements on ice in bulk that are given in Table 204.

69. THERMAL ENERGY OF ICE-I

This section is devoted to the following types of data for the ordinary type of ice (ice-I): the isopiestic specific heat (c_p) , the enthalpy or heat content (H = E + pv); the "free energy at constant pressure" (G = H - ST); and the entropy (S). No determination of either the heat of isothermal compression or the Joule-Thomson coefficient has been found for ice.

$$\int_{t_0}^{t} c_p dt = H_t - H_{t_0}; -T \int_{T_0}^{T} (H/T^2) dT = G_T - G_{T_0};$$

$$\int_{t_0}^{t} (c_p/T) dt = S_t - S_{t_0}.$$

Specific Heat of Ice.

In order to obtain the true specific heat of ice, proper allowance must be made for the progressive melting caused by the presence of included water containing dissolved impurities. Such inclusion is always present and gives rise to an apparent specific heat which exceeds the true, the excess increasing rapidly as $0\,^{\circ}$ C is approached. Even with the purest water used by Dickinson and Osborne, its effect was appreciable at $-5\,^{\circ}$ C, and became very marked above $-0.5\,^{\circ}$ C. Every nominal determination of the true specific heat of ice is to be regarded with suspicion unless the observer has clearly shown that this effect is negligible in his case, or has properly corrected for it. As early as 1904, A. D. Bogojawlensky 331a had concluded that the specific heat of a pure crystal is linear in the temperature.

Here we shall distinguish between the apparent specific heat (c_a) of ice and the true specific heat (c) which would be found were the ice perfectly pure. Obviously, the former will vary with the specimen, and the latter must be derived from the former. Apparently the only observers who have attempted to derive c from their own determinations are Dickinson and Osborne, who used four samples of very carefully purified water, and carried out the work with a precision unattained as yet by others. Over the range covered $(-0.5 \text{ to } -40 \,^{\circ}\text{C})$ they found that their results can be expressed by the formula $c_a = a + bt - d/t^2$ in which d varies with the specimen, but a and b do not. Hence, they regarded a + bt as the value

See Person, C. C., Ann. de chim. et phys. (3), 30, 73-81 (1850); Buchanan, J. Y.²⁸²; Smith,
 A. W., Phys. Rev. (2), 17, 193-232 (1903); Dickinson, H. C., and Osborne, N. S.²⁰⁰
 Bogojawlensky, A. D., Schrift, Natur. Ges. Univ. Jurjeff (Dorpat), 1904.

of c, and interpreted d/L, which decreases rapidly as the purity is increased. as the initial freezing-point of the solution obtained when the specimen is completely melted, L being the latent heat of fusion of ice at 0 °C. For the specimens used by them, d/L = -0.00125, -0.00120, -0.00095, and -0.00005 °C, respectively.

The best extended series of observations covering the range -2.9 to -189.5 °C is probably that of Nernst and his associates. 332 They represent their data by the formula $c_a = a + bt - d/t$, in which the last term varies as 1/t, whereas that in the Dickinson and Osborne formula varies as $1/t^2$. Their precision was not as high as that of Dickinson and Osborne.

Still lower temperatures (-189 to -250.6 °C) are covered by the observations of F. Pollitzer. 333 At such low temperatures $c_a = c$ except for experimental errors.

These and earlier observations were considered by J. H. Awbery in the derivation of the two values given for ice in his compilation.⁸³⁴

A second set of values is given by W. H. and E. K. Rodebush. 335 They have been computed by means of a formula, apparently unpublished, that was fitted to the observations of F. Pollitzer³³³ and of Nernst (as quoted by Pollitzer) and made c = 0 at 0 °K.

More recent determinations of c_a have been published by O. Maass and L. J. Waldbauer 336 and by W. H. Barnes and O. Maass. 337 These values are exactly those quoted by II. T. Barnes 338 and credited by him to "Maass and Barnes, W. H., 1927." The first covered the range -3 to -182.7 °C; the second, involving refinements in the method, covered the range -2.6 to -78.6 °C. In each case, the quantity measured was the total heat required to convert ice at $-t_1$ °C to water at $+t_2$ °C; t_2 was 16.5 °C in the first and 25 °C in the second. The quantity actually measured much exceeded the amount of heat accounted for by the specific heat of ice; consequently the precision with which the specific heat and its variation can be determined from those data is much lower than the precision of the data themselves. Only the latter precision is stated, which in the second article is said to be ±0.05 per cent or better. Actually, that is merely the precision of reproducibility for the same specimen under nominally identical conditions. No data are given from which any other precision can be determined. In each case, only a single specimen of ice seems to have been used.

The total heat (H) required to change the specimen from ice at t °C to water at 16.5 °C (25 °C in the second) was represented by a formula of the type $H = a + bt + dt^2 + et^3$ and c_a was obtained by differentiating that equation, giving $c_a = -b - 2dt - 3et^2$. Although no great precau-

 ³⁸² Nernst, W., Koref, F., and Lindemann, F. A., Sits. Preus. Akad. Wiss., 1910, 247-261 (1910); Nernst, W., Idem, 1910, 262-282 (1910).
 ³⁸³ Pollitzer, F., Z. Elektroch., 19, 513-518 (1913).

⁸⁸⁴ Awbery, J. H., Int. Crit. Tables, 5, 95-105 (95) (1929).

⁸⁸⁵ Rodebush, W. H., and E. K., Idem, 5, 89 (1929).

⁸⁸⁶ Maass, O., and Waldbauer, L. J., J. Am. Chem. Soc., 47, 1-9 (1925).

⁸⁸⁷ Barnes, W. H., and Maass, O., Can. J. Res., 3, 205-213 (1930). 888 Barnes, H. T., "Ice Engineering," p. 38, 1928.

tion was taken to ensure the purity of the water (ordinary distilled water was redistilled from an all-platinum still) there is nothing in the formulations to indicate any excessive increase in c_a as t approaches 0 °C, but those observations lying not below -110 °C in the first series can be satisfactorily represented by a formula of the type used by Dickinson and Osborne, d/L being taken as -0.0045 °C. The observers were of the opinion that at temperatures not exceeding -2.6 °C the ice was completely frozen, and consequently $c_a = c$. That opinion is not consistent with the observations either of Dickinson and Osborne or of Nernst, who used the best conductivity water. Actually, the precision with which they have determined the small quantities from which c_a and its variation have to be derived is much lower than they seem to have realized. It may be very conservatively taken as Δ/h where Δ is the excess of the observed value of H over that defined by their formula, and h is the excess of the observed value of H over that at 0 °C as defined by the formula; in accordance with their point of view, h is the heat required to raise the ice to 0 °C without melting it; the values of h, and they alone, are involved in the determination of c_a and its variation. All their values of h and of Δ are given below; it will be noticed that for no observation at a temperature exceeding -11 °C is Δ/h less than 1 per cent of h.

Maass and Waldbauer				Barnes and Maass					
t	H_{obs}	h	Δ	Δ/h	t	H_{obs}	h	Δ	Δ/h
-3.18	97.50	1.66	+0.12	+7.2%	-2.60	105.69	1.24	-0.03	-2.4%
-28.6	109.37	13.53	+0.08	$+0.6^{'}$	-4.60	106.73	2.28	+0.05	+2.2
-58.6	122.06	26.22	-0.11	-0.4	-10.15	109.40	4.95	+0.05	+1.0
-78.6	129.86	34.02	-0.24	-0.7	-15.17	111.75	7.30	+0.03	+0.4
-110.0	141.25	45.41	+0.23	+0.5	-19.99	113.95	9.50	-0.03	-0.3
-138.7	149.45	53.62	+0.34	+0.6	-25.01	116.31	11.86	+0.01	+0.1
-182.7	158.16	62.32	+0.08	+0.1	-30.05	118.69	14.24	+0.11	+0.8
					-50.05	127,28	22.83	+0.02	+0.1
					-78.57	138.21	33.76	-0.04	-0.1

In Table 205 the values of c_a as defined by the two formulas are compared. One might expect that the somewhat lower accuracy of the earlier work would be in large part offset by the much greater range of temperature over which the observations are spread, but the two formulas lead to markedly divergent results. Even at $-80\,^{\circ}\text{C}$ they differ by 7 per cent, the more recent giving the lower value. The observers' suggestion that a contributing factor to such divergence is the fact that the sublimation point of solid CO₂ was taken as $-78.2\,^{\circ}\text{C}$ in the earlier paper instead of as $-78.5\,^{\circ}\text{C}$ is not satisfactory, because according to their own formulation that change in temperature will change c_a by less than 0.2 per cent, which is negligible as compared with 7 per cent.

Until such marked discrepancies shall have been satisfactorily explained, confidence cannot be placed in either formula. This is especially unfortunate because the earlier work was in part intended to serve as a check upon the accuracy of similar measurements made upon other substances.

Table 205.—Apparent Isopiestic Specific Heat of Ice

The apparent specific heat of ice (c_a) exceeds the true (c) on account of the presence of impurities (see text); it varies from specimen to specimen. The best determinations are those of Dickinson and Osborne (DO), who found that $c_a = 0.5057 + 0.001863t - 79.75d/t^2$ cal₂₀/g·°C, where d = initial freezing point of the completely fused ice. Their values of d varied from -0.00005 to -0.00125 °C; if the impurity were NaCl and d were -0.000062 the concentration would be 0.0001 per cent by weight. Their observations did not extend below -40 °C.

Nernst and his associates extended their observations (N) to -189 °C and represented them by the formula $c_a = 0.470_2 + 0.0153_2t - 0.77_7/t$.

Pollitzer's observations (P) covered the range -189 to -250.6 °C.

The observations of Maass and Waldbauer (MW), and of Barnes and Maass (BM) were represented, respectively, by $c_a = 0.485 + 0.000914t - 5.46(10^6)t^2$ and $c_a = 0.48733 + 0.0009325t - 9.828(10^{-6})t^2$. For reasons stated in the text, implicit confidence cannot be placed in those values.

Excepting Pollitzer's, the values in Section 1 have been computed from these 4 equations, those lying beyond the range of the observations being enclosed in parentheses; $\delta = (BM - MW)/MW$.

The values of C in Section II have been taken directly from the paper cited; conversion has been made by the compiler.

Unit of $c_0 = 1$ calm per g.°C = 4.181 joules per g.°C; of $\delta = 1\%$.

-	~ ~ .		
	Various	determ	inations.

$\begin{array}{c} \operatorname{Ref}^{a} \to \\ d_{1} \to \end{array}$	-0 006	DO b	- 0.00006	N, P	MW	BM	
0 -5 -10 -20 -30	0.5155 0.4919 0.4696 0.4503	0.4983 0.4876 0.4685 0.4498	0.4966 0.4871 0.4684 0.4498	N 0.6180 0.5326 0.4784 0.4501	0.485 0.480 0.475 0.464 0.453	0.487 ₈ 0.482 ₄ 0.477 ₀ 0.464 ₈ 0.450 ₅	$ \begin{array}{r} \delta \\ +0.4 \\ +0.4 \\ +0.4 \\ +0.2 \\ -0.7 \end{array} $
-40 -60 -80 -100 -120	0.4315 (0.3940) (0.3568) (0.3194)	0.4312 (0.3939) (0.3567) (0.3194) (0.2821)	0.4312 (0.3939) (0.3567) (0.3194)	0.4283 0.3913 0.3571 0.3248 0.2929	0.440 0.410 0.377 0.339 0.297	0.434 ₃ 0.396 ₀ 0.349 ₈ (0.295 ₈) (0.233 ₉)	-1.4 -3.4 -7.1 -12.7 -21.2
-140 -160 -180		(0.2449) (0.2076) (0.1704)		0.2612 0.2299 0.1987	0.250 0.199 0.144	(0.164_2) (0.086_8) (0.001_0)	
-189 -200 -213.7 -222.3		(0.1331)		P 0.186 0.130 0.110			
-230.1 -235.4 -239.2 -244.8 -248.7 -250.6				0.098 0.085 0.070 ₀ 0.048 ₆ 0.037 ₀ 0.030 ₈			

Table 205—(Continued)

II. W. F. Giauque and J. W. Stout. 339 Mole fraction soluble impurity estimated to be 3×10^{-6} . Unit of C = 1 cal/g-mole. K; molecular weight taken as 18.0156; 1 cal = 4.1832 Int. joules.

$0 ^{\circ}\text{C} = 273.1 ^{\circ}\text{K}.$	Temp. = $t ^{\circ}C = T ^{\circ}$	K. Unit of $c_a = 1$	cal/g.
20 - 30 - 40 -	t -263.1 -253.1 -243.1 -233.1 -223.1	C 0.066° 0.490 0.984 1.466 1.896	0.0037 0.0272 0.0546 0.0814 0.1052
70 - 80 - 90 -	-213.1 -203.1 -193.1 -183.1 -173.1	2.304 2.701 3.075 3.448 3.796	0.1279 0.1499 0.1707 0.1914 0.2107
120 130 140	-163.1 -153.1 -143.1 -133.1 -123.1	4.130 4.434 4.728 4.993 5.265	0.2292 0.2461 0.2624 0.2772 0.2922
170 - 180 190	-113.1 -103.1 -93.1 -83.1 -73.1	5.550 5.845 6.142 6.438 6.744	0.3081 0.3244 0.3409 0.3574 0.3743
220 230 240	-63.1 -53.1 -43.1 -33.1 -23.1	7.073 7.391 7.701 8.103 8.326	0.3926 0.4103 0.4275 0.4448 0.4622
260 270	-13.1 -3.1	8.64 <i>2</i> 8.960	0.4797 0.4974

" References:

BM

DO

MW

Barnes, W. II., and Maass, O.³³⁷
Dickinson, H. C., and Osborne, N. S.³⁰⁰
Maass, O., and Waldbauer, L. J.³³⁰
Mernst, W., Koref, F., and Lindemann, F. A.,⁸³² and Nernst, W.³³²
Pollitzer, F.³³³

From the behavior of gadolinium anthraquinone sulfonate to which much water had been added, D. P. MacDougall and W. F. Giauque 340 have inferred that between 0.2 and 4 °K the specific heat of ice does not exceed 0.01 cal/g-mole °K.

^b For these values, 1 cal₂₀ = 4.183 joules, the value used by the authors in translating their electrical measurements into what they call cal 20.

This value for 10 °K has been derived by Giauque and Stout from the following values obtained in 1923 by Simon and privately communicated to them:

⁸⁸⁹ Giauque, W. F., and Stout, J. W., J. Am. Chem. Soc., 58, 1144-1150 (1936).

⁸⁴⁰ MacDougall, D. P., and Giauque, W. F., J. Am. Chem. Soc., 58, 1032-1037 (1936).

Table 206.—True Isopiestic Specific Heat of Ice

For temperatures not below -40 °C the best determinations are those of Dickinson and Osborne, which may be expressed by the formula c =0.5057 + 0.001863t cal₂₀ per $(g \cdot ^{\circ}C) = 2.115_3 + 0.00779_3t$ joule per $(g^{\circ}C)$. (They used the relation 1 cal₂₀ = 4.183 j.) Values so computed are given in column DO, those beyond the range of the observations being in parentheses.

Values observed by Nernst and by Pollitzer are given in column N, P; those at the higher temperatures are actually c_a , not c.

The two sets of values from International Critical Tables are given under A and R.

Ref ^a → DO A R 0 2.115 2.06 ± 0.01 2.1 ₂ -10 2.037 -20 1.959 1.94 ± 0.01 -23.1 1.935 1.9 ₃	N,P N 2.28 2.00
	2.28 2.00
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	2.00
-23.1 1.935 1.9 ₃	
	1 56
	1 56
-3 0 1.882	1 56
-40 1.804 1.82 \pm 0.01	1 56
$-60 \qquad (1.648) \qquad 1.68 \pm 0.02$	
$ \begin{array}{cccc} -70 & (1.570) \\ -73.1 & (1.546) & 1.32 \end{array} $	1.50
$ \begin{array}{lll} -73.1 & (1.546) & 1.32 \\ -80 & (1.492) & 1.54 = 0.02 \end{array} $	
$ \begin{array}{ll} -100 & (1.336) & 1.39 \pm 0.01 \\ -120 & (1.180) \end{array} $	1.23
-120 (1.180) -123.1 (1.152) 1.21	1,20
-140 (1.025)	
-150 (0.946) 1.030 ± 0.010	
-160 (0.868)	
$-170 \qquad (0.790)$	0.89_{6}
-173.1 (0.766) 0.8 ₀	_
-180 (0.712)	P
-189 (0.642)	0.77_{8}
$-200 \qquad (0.556) \qquad 0.653 \pm 0.013$	
-213.7 (0.450)	0.54
$-222.3 \qquad (0.383)$	0.462
-223.1 (0.377) 0.4 ₈ -230.1 (0.321)	0.411
	0.357
-235.4 (0.280)	0.337
-239,2 (0.251) -244.8 (0.207)	0.204
-248.7 (0.207)	0.15
-250 (0.167) 0.151 ± 0.004	
-250.6 (0.162)	0.128
-273.1 (-0.013)	

a References:

Compilation by Awhery, J. II. 384 based on the observations of Armstrong, H. E., Proc. Roy. Inst. Grt. Brit., 19, 354-412 (1908), Barnes, H. T., Trans. Roy. Soc. Canada, III (3), 3, 3-27 (1909), Dickinson, H. C., and Osborne, N. S., 300 Jackson, F. G., J. Am. Chem. Soc., 34, 1470-1480 (1912), Nernst, W., 382 and Ann. d. Physik (4), 36, 395-439 (1911), Nernst, W., Koref, F., and Lindemann, F. A., 382 Person, C. C., 281 Pollitzer, F., Z. Electrock., 17, 5-14 (1911); 19, 513-518 (1913), Regnault, V., Ann. d. Physik (Pogg.), 77, 99-109 (1849).

Dickinson, H. C., and Osborne, N. S. 300
Nernst, W., as quoted by Pollitzer, see P. Pollitzer, F. 388
Rodebush, W. H. and E. K., 385 see text.

DO

j/g.

Entropy of Ice.

If entropy is measured from 0 °K, then the entropy of water-vapor at 25 °C as computed from spectroscopic data is 45.10 cal/g-mole.°K 841; whereas the value found from the specific heat of ice and of water, together with the latent heats of transition, is only 44.28, a difference of 0.82 cal/ g-mole. K.342 Suggestions for explaining this discrepancy have been offered by W. F. Giauque and M. F. Ashley, 343 who definitely established the existence of the discrepancy, and by L. Pauling.³⁴⁴ Each suggestion attributes it to a failure of ice to attain the ideal state at the lowest temperature reached experimentally; the first assumes the persistence of the ortho and para molecular states at extremely low temperature, and the second assumes a certain amount of disorder. See also Table 207.

Table 207.—Various Isopiestic Thermal Data for Ice

 C_p = specific heat at constant pressure; $H_0 = \int_0^T C_p dT$ is the enthalpy (heat content); $G_0 = H_0 - ST = -\int_0^T (H_0/T^2) dt$ is the "free energy at constant pressure"; S_0 = entropy. All are for a pressure of 1 atm, and those with subscript o are measured from ice at $0 \, {}^{\circ}$ K; i =for ice, w =for water. For more complete definitions, see Table 1 and Section 6. values accepted by K. K. Kelley 354 essentially agree with the corresponding ones here given under R and S. 1 cal = 4.185 joules; 1 j/gfw = 0.05551

Unit of C_0 and $S_0 = 1$ i/(σ fw.°K): of H_0 and $G_0 = 1$ ki/ σ fw. Temp. = T°K: 1 σ fw = 18.0154g.

				kj/gtw. 1emp.		
$Ref^{a} \rightarrow T$	R	S	M	R	II	M
0	0	0	0	0	0	0
10	Ü	0.28	0.1	·	0.0006	0.0002
20		1.77	1.8		0.010	0.007
40	•	6.57	6.6		0.093	0.092
50	8.6		8.5	0.183		0.168
60		10.01	10.2		0.260	0.262
80		12.9	13.3		0.491	0.498
100	16.1	15.7	16.1	0.823	0.778	0.795
120			18.4			1.141
150	21.7	21.6	22.0	1.775	1.712	1.74
170			24.2			2.21
200	23.8	27.5	27.9	3.005	2.94	2.99
220			30.5			3.58
250	34.7	34.9	35.4	4.54	4.48	4.56
273.1 i	38.2	41.0	50.2	5.38	5.35	5.49
273.1 w		76.0	<i>7</i> 5.4		11.36	11.50

⁸⁴¹ Gordon, A. R., J. Chem'l Phys., 2, 65-72 (1934).

842 Giauque, W. F., and Stout, J. W., J. Am. Chem. Soc., 58, 1144-1150 (1936).

848 Giauque, W. F., and Ashley, M. F., Phys. Rev. (2), 43, 81-82 (L) (1933).

Pauling, L., J. Am. Chem. Soc., 57, 2680-2684 (1935).
 Kelley, K. K., Bur. Mines (U. S.), Bulletin 350 (1932).

Table	207-	(Continued)
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Ref.a→ T	R	S	M	R	S S ₀	M
0 10 20 40 50	0.044	0 0.00022 0.0036 0.0416	0 0.00006 0.00226 0.0348 0.0750	4.78	0 0.085 0.69 3.36	0 0.025 0.46 3.18 4.87
60 80 100 120 150	0.508 1.345	0.1424 0.3094 0.541 1.391	0.132 0.294 0.526 0.821 1.39	13.31 20.8	6.69 9.99 13.20 20.69	6.57 9.90 13.21 16.35 20.8
170 200 220 250 273.1 i 273.1 w	2.545 4.120 4.940 4.940	2.60 4.16 5.00 5.00	1.82 2.60 3.19 4.17 5.02 5.02	27.7 34.6 37.8 65.5°	27.7 34.4 37.9 59.9	23.7 27.9 30.8 34.9 38.4 60.5

" References:

- Miething, H., Abh. deuts. Bunsen-Ges., No. 9 (1920), based upon the data of Politzer (See R) and of Nernst, W., Ann. d. Physik (4), 36, 395-439 (1911), steps of 10 °K.
- Rodebush, W. H. and E. K., Int. Crit. Tables, 5, 84-91 (89) (1929), based upon the data of Pollitzer, F^{833} ; they measure both II and G from the uncombined gases at 0 °K and 1 atm. That is, their values for H and G fall below those given for H_0 and G_0 by the heat of formation of ice at 0 °K and 1 atm, which they have taken as 282.6 kj/gjw.

 Simon, F., "Hand. d. Physik," (Geiger and Scheel), vol. 10, 363 (1926) based on his own previously unpublished observations.

THERMAL CONDUCTIVITY OF ICE AND OF SNOW

Single Crystals.

The thermal conductivity of a single crystal of ice has not been studied, but indirect evidence indicates that the conductivity along the optic axis exceeds that in a direction perpendicular thereto.346 The difference is probably small. More recently, J. M. Adams 347 has reported observations which he thinks suggest "that the polar character of the crystal extends to the mechanism of thermal conduction in it."

The only recorded numerical data bearing upon the subject seem to be the following, k_v and k_h denoting the conductivities perpendicular and parallel, respectively, to the planes of freezing.

	kr	kn
Forbes (1873)	9.32	8.90 milliwatt/cm·°C
Straneo (1897)	21.9	21.0 milliwatt/cm.°C

Those by Forbes are surprisingly small.

^b Apparently erroneous. ^c For 25 °C (298.1 °K).

³⁴⁶ Barratt, T., and Nettleton, H. R., Int. Crit. Tables, 5, 231 (1929). Based upon: Barnes, H. T., Nature, 83, 276 (1910), Forbes, G., Proc. Roy. Soc. Edinburgh, 8, 62-69 (1873), Straneo, P., Atti. Acc. Lincei (5), 6₂: 299-306 (1897).

⁸⁴⁷ Adams, J. M., Proc. Roy. Soc. London (A), 128, 588-591 (1930) → Phys. Rev. (2), 36, 788 (A) (1930).

Ice in Bulk.

When data were being prepared for the International Critical Tables, the most extended series available on the thermal conductivity of ice was that by C. H. Lees.³⁴⁸ More recent work by M. Jakob and S. Erk,³⁴⁹ extending to -130 °C, is probably to be preferred to all others now available. They found that when there is a flow of heat between a block of ice and a metal plate frozen to it, there is always a discontinuity in the temperature at the junction, the discontinuity increasing as the temperature is lowered.

Apparently, the earliest recorded attempt to measure the thermal conductivity of ice is that of F. Pfaff 350 leading to the surprising conclusion that the conductivity of ice is 0.82 that of iron (i.e., to k = 508 milliwatt/cm.°C).

Table 208.—Thermal Conductivity and Diffusivity of Ice

(For single crystals, see text.)

The preferred values for the conductivity (k) are those (JE^a) by Jakob and Erk. The (VDa) values have been computed by Van Dusen's equation $(k = 20.9 (1 - 0.0017t) \text{ milliwatt/cm} ^{\circ}\text{C})$, which was set up prior to the work by JE, and approximately represents the values found by Lees.

Thermal diffusivity $(k/\rho c, \rho = \text{density}, c = \text{specific heat})$ is 0.011 cm²/sec if t > -30 °C (VD^a); is 0.0114 at 0 °C (SH^a), based on F. Neumann. 351

Unit of	k=1	milliwatt/	(cm °C)
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Ref.a→						Ref.a→			Ref a→	\mathbf{v}_{D}
t		k	t		k	t		k	t	ĸ
0	22.48	20.9	-50	27.8	22.7	-100	34.7	24.4	-150	26.2
-10	23.2	21.3	-60	29.1	23.0	-110				26.6
-20	24.3	21.6	-70	30.5	23.4	-120	38.1	25.2	-170	26.9
-30	25.5	22.0	-80	31.8	23.7	-130	40.2	25.5		
-40	26.6	22.2	-90	33.1	24.1	-140		25.9		

a References:

Ces:

Jakob, M., and Erk, S. S49

Schofield, F. H., and Hall, J. A., Int. Crit. Tables, 2, 315-316 (1927); based on Ingersoll, L. R., and Zobel, O. J., "An Introduction to the Mathematical Theory of Heat Conduction," 1913; Neumann, F. S51 Straneo, P. S40

Van Dusen, M. S., Int. Crit. Tables, 5, 216-217 (1929); based on Lees, C. H. S48; Abels, H., Repert. f. Meteor. (Wild's, St. Petersburg), 15, No. 1 (1893); Andrews, T., Proc. Roy. Soc. (London), 40, 544-549 (1886); Forbes, G. S46; Mitchell, A. C., Proc. Roy. Soc. Edinburgh, 13, 592-596 (1886); Straneo, P., Nuovo Cim. (4), 7, 333-340 (1898) & Atti Accad. Lincei (5), 62: 262-269, 299-306 (1897). VD

^b S. Arzybyschew and I. Parfianowitsch ⁸⁵² find 23.0; P. G. Tait, ⁸⁵⁸ 21; and SII gives 22 mw/cm·°C.

³⁴⁸ Lees, C. H., Phil. Trans., 204, 433-466 (1905) → Proc. Roy. Soc. (London), 74, 337-338 (1905).

³⁴⁹ Jakob, M., and Erk, S., Wiss. Abh. d. P. T. R., 13, 395-409 (1929) = Z. gcs. Kälte-Ind., 36, 229-234 (1929) \rightarrow Z. techn. Physik, 10, 623-624 (1929).

⁸⁵⁰ Pfaff, F., Sitzb. physik.-mcd. Soc. Erlangen, 6, 155-157 (1874).

⁸⁵¹ Neumann, F., Phil. Mag. (4), 25, 63-65 (1863).

²⁶² Arzybyschew, S., and Parfianowitsch, I., Z. Physik, 56, 441-445 (1929).

²⁶⁸ Tait, P. G., Proc. Roy. Soc. Edinburgh, 13, 592-596 (1886).

Snow.

The thermal conductivity (k) and diffusivity (D) of snow vary greatly with the density, and the values obtained by different observers exhibit much discordance (see Table 209).

Table 209.—Thermal Conductivity and Diffusivity of Snow

The thermal conductivity (k) and diffusivity $(D = k/\rho c)$, both depend upon the density of the snow; ρ = density, c = specific heat. From a consideration of the data then available, M. S. van Dusen 358 concluded that between 0 and $-30 \,^{\circ}\text{C}$, $k = 0.21 + 4.2 \,\rho + 21.6 \,\rho^{3}$ milliwatt/(cm. $^{\circ}\text{C}$), $1000D = 2.0 + 0.1/\rho + 10.3 \rho^2$ cm²/sec, where ρ gm/cm³ = density of the snow (not of the individual crystals). More recently, J. Devaux 359 has concluded that $k = 0.29 (1 + 100 \rho^2)$ milliwatt/(cm.°C). Values computed by means of these equations are here entered under VD or D, as may be appropriate.

Unit of k = 1 milliwatt/(cm.°C) = 239 10-6 g-cal/(cm.sec.°C); of D = 1 cm².sec; of $\rho = 1$ g/cm³. Temp. = 0 °C

Ref.a→	VD	D		Ref.a	Ref.a→	VD	000 D S	H Ref.
0.11	0.70	0.65	1.07	J	0.125	3.08		Acci.
0.125	0.78	0.75			0.19	2.9	2.50	A
0.24	1.52	1.98	1.67	OAY	0.33	3.4	4.60	A
0.25	1.60	2.1	1.88	OAY	0.40	3.9		
0.27	1.8	2.4	1.34	OĄY	0.50	4.8	4.1	T 17.7
0.45 0.50	4.1 5.0	6.2 7.6	0.49 1.3	17	•		4.1	IK
0.50	J.U	7.0	1.0	14				

- References.
 - A
 - ίĸ
 - Abels, H., Rep. Metcor. (Wild, St. Petersburg), 16, No. 1 (1893).
 Devaux, J. Sto.
 Ingersoll, L. R., and Koepp, O. A., Phys. Rev. (2), 24, 92-93 (1924).
 Ingersoll, L. R., and Zohel, O. J., "An Introduction to the Mathematical Theory of Heat Conduction," 1913.
 - Jansson, M., Öfvers. af K. Svenska Vet. Akad., Förh. (Stockholm), 58, 207-222

 - OAY Okada, T., Abe, K., and Yamada, J., Proc. Tokyo Math.-Phys. Soc. (2), 4, 385-389 (1908).

 SII Values tabulated by Schofield, F. H., and Hall, J. A., Int. Crit. Tables, 2, 313-315 (1927) and ascribed to the reference here given in the adjacent column.

 VD Van Dusen, M. S. Sos.; equation based on observations of A, IK, J, OAY, Hjeltström, S. A., Öfvers. af K. Svenska Vet. Akad. Förh. (Stockholm), 46, 669-676 (1889), and Neumann, F., Ann. de chim. et phys. (3), 66, 183-187 (1862).
- ^b For $\rho = 0.125$, M. Kuroda ⁸⁶⁰ gives k = 2.16. 1000D = 8.6.

Snow densely packed.

From his work on the névé on the Mt. Blanc glaciers at elevations of 4.2 to 4.4 km, J. Vallot ³⁵⁴ found that the diurnal variation in temperature did not extend below one meter, at which depth the temperature rises to 0° C only under exceptional conditions; the annual variation does not extend below 6.5 m. He observed the following temperatures, t_1 in the year 1900, t_2 in 1911:

²⁵⁴ Vallot, J., Compt. rend., 156, 1575-1578 (1913).

M. Kuroda ³⁵⁵ observed that the lowest temperature in a snow-blanket 9 meters thick occurred at a depth of about 4.5 meters, and was -0.9 °C, the temperature of the surface being +0.2 °C, and of the ground +0.4 °C. The density varied from 0.35 g/cm⁸ at 50 cm to 0.65 at 7 meters.

The distribution of temperature in the snow-blanket and in the underlying soil, near Sodoakylä, Finland, within the Arctic Circle, has been

studied over a period of 24 months by J. Keränen. 356

The Oxford University's Arctic expedition, of 1935-6, to North-East Land found that at a depth of 70 ft. (21 m) in the ice cap the temperature was fairly constant at 0.0 °C; and at a somewhat greater depth an unfrozen lake was found.⁸⁵⁷

71. REFRACTIVITY OF ICE

Over a hundred years ago, Sir David Brewster 361 observed that crystals of ice are optically uniaxial and positive, the index of refraction (ϵ) of the extraordinary ray exceeding that (ω) of the ordinary. A. Bertin 362 has stated that the interference fringes seen when ice is suitably observed in a polarizing microscope are among the most beautiful exhibited by any uniaxial crystal.

H. E. Merwin ³⁶³ has concluded from the observations of A. Ehringhaus ³⁶⁴ that between -3 °C and -65 °C, and $405 \,\mathrm{m}\mu$ and $706 \,\mathrm{m}\mu$. $\epsilon_{\lambda} - \epsilon_{D} = 1.01 \,(\omega_{\lambda} - \omega_{D})$ and $d\omega/dt = -3.8 \times 10^{-5}$ per °C, the subscript denoting the wave-length to which the index applies, D denoting the D-lines of sodium at $\lambda = 589.3 \,\mathrm{m}\mu$. Hence, $d\epsilon/dt = 3.84 \times 10^{-5}$ per °C. From these relations and other data given in Merwin's compilation, the data in Table 210 have been derived. The form of the factors F_{ω} and F_{ε} was empirically determined for this compilation.

Taking for the density of ice at -3 °C the value 0.9164 g/cm³ (Tables 200 and 202), and for the indices of refraction for the D-lines the values given in Table 210, the Lorenz "refraction constant" $\left(\frac{n^2-1}{\rho(n^2+2)}\right)$ is 0.2097 cm³ per gram for the ordinary, and 0.2105 for the extraordinary ray. The corresponding quantity for water at 20 °C (n=1.33300, $\rho=0.9982$ g/cm³) is 0.2061, and for water-vapor at 1 atm and 110 °C ($n=1+313.30 \rho \times 10^{-6}$ (Table 58), $\rho=0.0005804$ g/cm³) is 0.2088 cm³/g. Early and rough determinations of ω and ε were made by E. Reusch, ³⁶⁵

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855 Kuroda, M., Sci. Papers Inst. Phys. and Chem. Res., Tokyo, 12, 69-81 (1929).
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³⁵⁶ Keränen, J., Annales. Acad. Sci. Fennicae (A), 13, No. 7 (1920).

as7 Glen, A. R., Nature, 139, 10-12 (1937).

⁸⁵⁸ Van Dusen, M. S., Int. Crit. Tables, 5, 216 (1929).

³⁵⁹ Devaux, J., Ann. de phys. (10), 20, 5-67 (1933).

²⁶⁰ Kuroda, M., Sci. Papers Inst. Phys. and Chem. Res., Tokyo, 12, 149-159 (1930).
²⁶¹ Brewster, Sir David, Phil. Trans., 1814, 187-218 (1814); 1818, 199-273 (1818); Phil. Mag. (3), 4, 245-246 (1834).

see Bertin, A., Ann. de chim. et phys. (3), 69, 87-96 (1863).

ses Merwin, H. E., Int. Crit. Tables, 7, 17 (1930).

⁸⁰⁴ Ehringhaus, A., Neues Jahrb. Mineral., Geol., Beilage Bd. 41, 342-419 (1917).

²⁶⁵ Reusch, E., Ann. d. Physik (Pogg.), 121, 573-578 (1864).

and fairly precise ones covering most of the visible spectrum, by C. Pul-frich. 866

Table 210.-Indices of Refraction of Ice

Adapted from compilation of H. E. Merwin.³⁶³ For long waves, see Tables 218 and 219.

 ω , ϵ = index of refraction of the ordinary and the extraordinary ray, respectively. Subscript indicates either the wave-length to which the index refers, D indicating the wave-length of the D-lines of sodium (= 5893A), or the Centigrade temperature. Between -3 and -65 °C and in the range λ = 4046A to 7065A the following relations hold good: $\omega_t = \omega_{-3} - (11.4 + 3.8t) \times 10^{-5}$; $\epsilon_t = \epsilon_{-3} - (11.5_2 + 3.84t) \times 10^{-5}$; $\epsilon_{\lambda} - \epsilon_D = 1.01(\omega_{\lambda} - \omega_D)$; approximately, $\omega_{\lambda} = \omega_D + (\lambda_D - \lambda)F_{\omega}$, $\epsilon_{\lambda} = \epsilon_D + (\lambda_D - \lambda)F_{\epsilon}$, where $10^6F_{\omega} = 2.07 + 10^{(\lambda_D - \lambda)/4000}$ per angstrom, $10^6F_{\epsilon} = 2.09 + 1.01 \times 10^{(\lambda_D - \lambda)/4000}$ per angstrom, all λ 's being expressed in angstrom units. The order of approximation may be seen by comparing the tabulated values of $10^4(\lambda_D - \lambda)F_{\omega}$ with those of $10^4(\omega_{\lambda} - \omega_D)$ derived from the values of ω ; interpolation is facilitated by the use of the values of the F's.

Unit	of $\lambda = 1A = 10^{-8}$	cm. Temp.	= -3 °C. I	ndex with reference	to a	vacuum
λ	(n)	€	$104(\omega_{\lambda}-\omega_{D})$	$10^4(\lambda p - \lambda)F_{\omega}$	10^6F_{ω}	$10^6 F_{\epsilon}$
4046 Hg	1.3183	1.3198	93	91.6	4.96	5.02
4358 Hg	1.3159	1.3174	69	69.0	4.49	4.53
4861 H	1.3129	1.3143	39	40.1	3.88	3.92
4916 Hg	1.3126 +	1.3140 +	36+	37.3	3.82	3.86
5461 Hg	1.3104	1.3118	14	14.5	3.35	3.38
5780 Hg	1.3093 +	1.3107	3+	3.5	3.14	3.17
5893 Na	1.3090	1.3104	0	0	3.07	3.10
6234 Hg	1.3079	1.3093	-11	-9.9	2.89	2.92
6563 H	1.3070 +	$1.3084 \pm$	-19+	-18.4	2.75	2.78
6908 Hg	1.3063	1.3077	-27	-26.7	2.63	2.65
7065 He	1.3060	1.3074	-30	-30.2	2.58	2.60

72. REFLECTIVITY OF ICE AND OF SNOW

Ice.

Throughout the visible spectrum the transparency of ice is so great that the reflectivity (R) can be computed satisfactorily from the index of refraction by means of the formulas given in Section 38. The observed reflectivity passes through a pronounced maximum at $\lambda = 3.2 \,\mu$ and again near $\lambda = 13.0 \,\mu$. M. Weingeroff ⁸⁶⁷ suggests that in the latter region there are "residual" rays.

Defining the reflectivity as $R = I_r/I_t$, I_t and I_r being the intensities of the incident and of the specularly reflected radiation, respectively,

⁸⁰⁶ Pulfrich, C., Idem (Wied.), 34, 326-340 (1888).

sor Weingeroff, M., Z. Physik, 70, 104-108 (1931).

E. P. T. Tyndall ³⁶⁸ has given the following values, based upon the observations of G. Bode, ³⁶⁹ angle of incidence about 15°.

$$\lambda$$
 1.0 1.5 2.0 2.4 2.6 2.8 3.0 3.2 3.4 3.5 4.0 μ 100 R 1.72 1.62 1.62 1.13 0.73 0.70 1.60 5.10 3.90 2.81 1.75

M. Weingeroff ³⁶⁷ has observed the following, the angle of incidence being about 12°:

$$\lambda$$
 6.0 7.5–9.0 10.0 10.5 11.0 11.2 11.5 12.0 12.5 13.0 13.8 14.3 15.0–16.0 μ 100R 0.8 0.5 0.4 0.5 1.0 1.5 2.0 2.5 3.0 3.5 3.0 2.5 2.0

It has been reported that the intensity of the reflection of x-rays from the (100) plane of ice is reduced about 2.5 per cent by an electric field of 1300 volts/cm parallel to that plane.³⁷⁰

Snow.

Freshly fallen, powdery snow on mountains is mat and closely obeys Lambert's law.³⁵⁹

Using filtered radiation from a quartz-enclosed mercury arc, and working at a vertical angle of 40° between the arc and the receiving instrument, E. O. Hulburt ³⁷¹ found the effective (diffuse) reflecting power (R_c) of freshly fallen snow to have the following relative values, that for the region $\lambda = 0.4$ to $0.8 \,\mu$ being arbitrarily taken as 100:

$$\lambda$$
 0.3 to 0.4 0.4 to 0.8 0.8 to 2.6 2.6 to 7 Beyond 7 μ 88 100 38 45 65

The albedo of a plane surface is defined as $A = F_r/F_t$, where $F_r =$ total luminous flux reflected by the surface when uniformly illuminated by white light, the total luminous flux incident on the surface being F_t . From the observations of P. G. Nutting, L. A. Jones, and F. A. Elliott, ³⁷² E. P. T. Tyndall ³⁷³ concluded that A = 0.93 for snow, A for MgCO₃ being assumed to be 0.98; J. Devaux ³⁵⁹ gives A = 0.95.

These values are much higher than those reported by others. From a long series of observations made near Leningrad, N. N. Kalitin 374 found the maximum value $\mathcal{A}=0.87$ for dazzling, fresh, soft snow fallen the evening before. He quotes the following previously reported values for such maximum:* C. Dorno, 0.89; Abbot and Aldrich, 0.70; A. Ångström, 0.81. The packing of the snow with age decreases the albedo; so does melting of the surface. He found the apparent albedo to lie above 0.45 so long as the ground was completely covered with snow that was not more than

^{*}Values ranging from 0.70 to 0.89 have been reported by H. H. Kimball and I. F. Hand. 874a

⁸⁰⁸ Tyndall, E. P. T., Int. Crit. Tables, 5, 256-263 (258) (1929).

Bode, G., Ann. d. Physik (4), 30, 326-336 (1909).

⁸⁷⁰ Német, A., Helv. Phys. Acta, 8, 97-116 (1935).

⁸⁷¹ Hulburt, E. O., J. Opt. Soc. Amer., 17, 23-25 (1928).

⁸⁷⁸ Nutting, P. G., Jones, L. A., and Elliott, F. A., Trans. Illum. Eng. Soc. (N. Y.), 9, 593-597 (1914).

ers Tyndall, E. P. T., Int. Crit. Tables, 5, 262 (1929).

⁸⁷⁴ Kalitin, N. N., Monthly Weather Rev., 58, 59-61 (1930).

⁸⁷⁴a Kimball, H. H., Hand, I. F., Monthly Weather Rev., 58, 280-281 (1930).

slightly soiled. The following are typical of the values he obtained 375: Loose surface, 0.80; freshly fallen snow, 0.83; dense surface, 0.86; thawed and grainy surface, 0.40; uneven surface, 0.75.

73. LUMINESCENCE OF ICE

By the luminescence of a substance is meant its emission of light under the existing conditions, and in particular from its interior, as distinguished from reflection by its surface. Several types of luminescence are described in Section 39. For the internal brilliance of a blanket of snow, see Section 75. For reports of the crystalloluminescence of ice, see end of Section 97.

Fluorescence of Ice.

Under this head, phosphorescence, triboluminescence, etc, are included. While exposed to the filtered radiation from radium, the filter being Pt 2 mm thick, ice fluoresces, but less brightly than does water at 20° and under the same conditions. In both cases the luminescence is very weak.³⁷⁶

Rayleigh Scattering by Ice.

The vivid blue color of large masses of pure ice has been ascribed by C. V. Raman to the scattering of light by the molecules of the ice, or rather by the slight variations in the concentration of the molecules (cf. Section 39). The purer the ice, the deeper the blue. Slight traces of impurities alter the color very perceptibly.377

Raman Scattering by Ice.

For an account of the general characteristics of the Ramah effect, see Section 39, and K. W. F. Kohlrausch. 378 Each Raman band for ice is much narrower than the corresponding one for water and corresponds to a slightly smaller value of δv , the difference between the wave-number of the Raman band and that of its exciter.³⁷⁹ Early observations by I. R. Rao ³⁸⁰ indicated that the intensity of the unresolved prominent band for ice depends upon the frequency of the exciter. He reported as follows, λ_{Hg} being the wave-length of the exciter, and I the intensity of the corresponding Raman-band: $\lambda_{Hg} = 3650A$, I = 15; $\lambda_{Hg} = 4047A$, I = 10; $\lambda_{Hg} =$ 5060A, I = 5. No later observation on such variation has been found.

Effect of temperature.—At the temperature of liquid air (ca. - 190 °C) the Raman spectrum of ice consists of one intense and fairly sharp line at $\delta \nu = 3090$ cm⁻¹ ($\lambda_R = 3.24 \,\mu$) and a faint companion at $\delta \nu = 3135$ cm⁻¹ $(\lambda_R = 3.19 \,\mu)$; whereas at temperatures near 0 °C it consists of diffuse lines or bands at $\delta \nu = 3196$ cm⁻¹ ($\lambda_R = 3.13 \,\mu$) and 3321 cm⁻¹ ($\lambda_R =$

⁸⁷⁵ Kalitin, N. N., Gerlands Beitrag. s. Geophysik, 34, (Köppen Bd. 3), 354-366 (1931).

Mallet, L., Compt. rend., 183, 274-275 (1926).
 Barnes, H. T., "Ice Engineering," pp. 8, 9, 1928.
 Kohlrausch, K. W. F., "Der Smekal-Raman Effekt," 1931.
 Ganesan, A. S., and Venkateswaran, S., Indian J. Phys., 4, 195-280 (1929).

⁸⁵⁰ Rao, I. R., Idem, 3, 123-129 (1928).

3.01 μ . For interpretations of the change, see also I. R. Rao, 382 who disagrees with Sutherland.

Table 211.—The Raman Spectrum of Ice

 δv is the difference between the wave-number $(1/\lambda)$ of the Raman line and that of the exciting radiation; $\lambda_R = 1/\delta_V$. Each number or check mark in the columns of relative intensity is placed on the line with the value of δν that corresponds to the maximum of the line or band as reported by the indicated author. The absence of such number or mark indicates that the author did not report a maximum at that value of δv .

		Unit of $\delta v = 1$	cm ⁻¹ ; o	of $\lambda_R = 1 \mu$	= 10-4cm		
	Ref.ª→	Rao 1928	GaV 1929	Ras 1932	Rao 1934	CBL 1937	Hib 1937
$\delta \nu$	λ_R			- Relative i			
53.5	187			1d			
205	48.8						4
210	47.6					\checkmark	
212.1	47.15			5d			
601	16.6						3
2225	4.494						2 10
3136	3.189						10
3150	3.175					√c	
3156	3.168					√c ✓	
3190	3.135	5				•	
3193	3.132		v /				
3196	3.129		V		55		
3200	3.125	10					
3270	3.058	15					
3300	3.030					√c	
3321	3.011			•	40	•	
3330	3.003				10		8
3390	2.950		1/	i		√c	U
3420	2.924		v	1		VC	
3549	2.818		1/	•			
5393	1.854		V				

Rao 382 has given the following values for the relative intensity of the Raman scattered light throughout the range $\delta_{\nu} = 2877$ to 3768 cm⁻¹.

$\stackrel{oldsymbol{\delta u}}{I}$	2877 0	3019 10	3122 36	3196 55	3252 37	3321 40
δν	3321	3389	3466	3538	3636	3768
1	40	.39	.30	12	5	0

a References:

Cross, P. C., Burnham, J., and Leighton, P. A., J. Am. Chem. Soc., 59, 1134-1147 (1937).

Rasetti reported a continuous band extending from $\delta v = 3300$ to $\delta v = 3420$ cm⁻¹.

GaV

Ganesan, A. S., and Venkateswaran, S. ³⁷⁸
Hibben, J. H., J. Chem'l Phys., **5**, 166-172 (1937).
Rao, I. R., 1928, ³⁸⁰ 1934. ³⁸⁹
Rasetti, F., Nuovo. Cim. (N. S.), **9**, 72-75 (1932).

^b The numerical values of the relative intensities have no significance except with reference to others appearing in the same column. When a reference contains no numerical estimate of the relative intensities of the lines or bands recorded, the positions of those lines or bands are indicated by a check mark (\checkmark) ; d indicates that the band was recorded as being diffuse; c indicates the value of $\delta \nu$ corresponding to a fundamental frequency of a band. The long line in Ras column indicates that

74. DIFFRACTION OF X-RAYS BY ICE

The diffraction of x-rays by ice has been studied primarily for the purpose of ascertaining the intimate crystalline structure of ice, and most of the reports of such work contain no explicit statement of the values of the individual periodicities observed, or of their relative intensities. The information obtained regarding the crystalline structure is given in Section 60, on the molecular data for ice. Values of the observed periodicities and their relative intensities are given in Table 212.

Table 212.—Diffraction of X-rays by Ice

 $d=(\lambda/2)\cdot\sin{(\phi/2)}$, $\lambda=$ wave-length of the incident x-rays, $\phi=$ angle of diffraction at which the intensity of the diffracted radiation passes through a maximum; d characterizes some kind of periodicity in the structure of the crystal. I= relative intensities of the several maxima; w= weak, s= strong, m= medium strong, v= very; e.g., vvs= very, very strong. Unit of d=1A.

•	,									
						B() a				
StJ 4	Det	n a	- 50 to	- 80°	- 8	5 °C	- 90 to	105° -	- 115 to -	- 175°
d	d	I	d	I	d	I	d	I	ď	I
4.15										
	3.92	10	3.90	w	3.87	w	3.90	w	2 -	
	3.67	100	3.63	vvs	3.69	vvs	3.66	w s	3.7	m
3.46	3.44	20	3.40	m	3.42	m	0.00	5	,	
3.30	0.11	20	0.40	111	0.72	311				
0.00	2.68	15	2.64	***	2.66	***				
2.56	2.00	13	2.04	m	2.00	m				
2.34		4.0						_		
	2.26	10	2.26	w	2.28	vw	2.25	S)	
	2.065	50	2.05	vs	2.05	vs		S	2.1	w
1.94	1.92	10	1.90	w	1.92	w	1.91)	
			1.71	w		••				
	1.516	15	1.51	w	1.52	w				
	2.020		1.45	m	1.02	**				
	1.368	20	1.35	m	1.37	m				
4			1.55	111	1.5/	111				
1.30	1.30	2.5								
1.26	1.25	2.5	1.26	vw						
			1.21	vw						
	1.167	5	1.17	vw	1.17	vw				
0.74										

a References:

BO Burton, E. F., and Oliver, W. F., Proc. Roy. Soc. (London) (A), 153, 166-172 (1936).

Den Dennison, D. M., Phys. Rev. (2), 17, 20-22 (1921).

St. John, A., Proc. Nat. Acad. Sci., 4, 193-197 (1918).

75. Absorption and Transmission of Radiation by Ice and by Snow

(There seem to be no such data for x-rays, γ -rays, cosmic radiations, or corpuscular radiation. For $\lambda > 4 \mu$, see Section 80; for scattering by ice, see Section 73.)

⁸⁸¹ Sutherland, G. B. B. M., Proc. Roy. Soc. (London) (A), 141, 535-549 (1933).

⁸⁸² Rao, I. R., Idem, 145, 489-508 (1934).

Table 213.—Monochromatic Absorptivity of Ice

Trans. = per cent transmitted. The absorptivity (k) is defined by the equation $I = I_0 e^{-kx}$, where $(I_0 - I)$ is the decrease in the intensity caused by transmission through x cm of ice in the interior of the block.

Unit of
$$\lambda = 1 \mu = 10^{-4}$$
 cm; of $x = 1$ cm; of $k = 1$ cm⁻¹

I. Plane-polarized radiation transmitted perpendicular to the optic axis; x = 0.5 cm; $\lambda =$ wave-length at which the absorptivity passes through a maximum. P.a

	Ordinary ray	b	Ex	traordin <mark>ar</mark> y ra	у в
λ	Trans.	\boldsymbol{k}	λ	Trans.	k
0.79	93	0.145	0.81	55	1.20
0.89	93.5	0.137	0.92	54	1.23
1.02	93	0.145	1.06	46	1.55
1.26	57	1.12	1.29	27	2.62

Unpolarized radiation transmitted probably parallel to optic axis.

III. K^a Crystallographic direction is not stated. The reported transmissions by the two blocks are not consistent. If they are combined on the assumption that the blocks differ solely in the value of x, then one finds for the reflectivity (R) a negative value (-5%) and for k the values here given. If R is assumed to be negligible, and only the data for the longer block are used, one obtains the values k_{107} .

λ	0.332	0.346	0.366	0.392	0.416	0.438	0.446
Trans., $x = 10$	97	96	99	99	98	99	98
Trans., $x = 107$	46	46	51	52	54	52	55
10'k	77	76	68	66	62	65	59
10^4k_{107}	73	73	63	61	58	61	56

IV. CE^a Crystallographic direction is not stated. Values of nk are reported for the residual rays from the salts indicated. Thickness of ice was x = 0.0034 cm. Temp. = -10 °C.

Salt	NaCl	KCl	KBr	TICI	TlBr	TII
λ	52	63	83	100	117	152
Trans.	11	13	32	72	85	84
$\lambda k/4\pi$	0.27	0.30	0.22	0.08	0.03	0.03
k	650	600	330	126	30	25

a References:

Dode, U. 5000 Cartwright, C. H., and Errera, J., Proc. Roy. Soc. (London) (A), 154, 138-157 (1936)—Acta Physicochim. URSS, 3, 649-684 (1935) → Cartwright, C. H., Nature, 136, 181 (L) (1935). Kalitin, N. N., Compt. rend. Acad. Sci. URSS (N. S.), 9 = 19354, 145-146 (1935). Plyler, E. K.384

b Certain inconsistencies in this and his other paper and reporting the same data have resulted in several quotations in which the data for the ordinary ray have been assigned to the extraordinary, and conversely. The assignment here given is that in the author's Table I, which he has informed me is correct.

The absorption is given as 93 per cent, which leads to k = 26.7 cm⁻¹, a value far

exceeding what one would infer from Plyler's data.

For a given layer of material, the transmitted fraction of the incident radiation depends upon the amount of the radiation that is scattered by the layer, as well as upon the amount that is truly absorbed, that is converted into another form of energy. But this distinction has not been observed in the reporting of experimental data for the absorption of ice, the entire reduction in intensity being described as absorption. The error so produced is probably very small when the ice is clear, except for the shorter wave-lengths, for which measurements of the absorption seem to be lacking. For snow, the scattering is of prime importance; the true absorption is that of the individual ice-crystals, of the ice itself.

Ice.

In the visible spectrum, the absorptivity of ice is certainly small. In the infrared, beyond $\lambda=1~\mu$, it is great and entirely analogous to that of water (Section 43), at least as far as $\lambda=6~\mu$. A plate of ice 1 mm thick absorbs practically all radiation for which $\lambda \ge 3~\mu$, and a frozen soap-film cuts off nearly all radiation for which $\lambda \ge 6~\mu$. For λ greater than about $4~\mu$, the absorptivity of ice is great.^{259, 383}

The extraordinary ray is more strongly absorbed than the ordinary, and the corresponding wave-lengths at which the maxima of the absorption occur are greater in the former than in the latter.³⁸⁴

The intercrystallic material, which in some cases was found to be less than 0.0008 cm thick, "has a much higher absorption of infrared light than the ice itself." ³⁸⁵ Plyler concluded that this extra absorption was not due to dissolved salts.

Table 214.—Transmissivity of Ice for Black-Body Radiation

Adapted from data given by S. L. Brown.³⁸⁷

Thickness of ice = 3 mm.^a Transmission = τ per cent; τ_c is defined by an empirical equation ($\tau_c = -18.5 + 0.033t$) constructed for the present compilation. Temperature of the source of radiation is t °C.

t	τ	τς	t	T	τ_o
660	3 4	3.3	865	9.9	10.0
720	5.2	5.3	910	11.7	11.5
790	7.6	7.6	925	11.9	12.0
			960	14.2	13.2

^a H. Hess ⁸⁸⁸ has stated that Melloni found that a plate of ice 2.6 mm thick transmitted only 6 per cent of the total radiation incident on it from a Locatelli lamp.

Snow.

In a study of the penetration of radiation into snow and glaciers the

Angström, A., Ark. f. Math. Astr., och Fysik, 13, No. 21 (1919).
 Schaefer, C., and Matossi, F., "Das ultrarote Spektrum," 1930; Plyler, E. K., J. Opt. Soc. Amer., 9, 545-555 (1924).

³⁸⁸ Plyler, E. K., J. Elisha Mitchell Soc., 41, 18 (1925).

⁸⁸⁶ Plyler, E. K., J. Elisha Mitchell Soc., 41, 39-40 (1925).

²⁸⁷ Brown, S. L., Phys. Rev. (2), 21, 103-106 (1923).

²⁰⁶ Hess, H., "Die Gletscher," 1904.

following terms are useful: Factor of entrapment (E) = unity minus the albedo (Section 72) = fractional excess of radiation incident upon the surface over that returned by the snow or glacier. Internal illumination (I_i) = sum of the flux of radiation each way through a given unit surface at the place considered; it may be expected to vary with the aspect of the surface. Factor of attenuation (T) = ratio of internal illumination to the illumination (I_g) of the surface of the snow or glacier; $T = I_i/I_g$.

Clean, freshly fallen snow has a perfectly diffusing surface, the light proceeding from it when illuminated being distributed in accordance with Lambert's law,* and entirely independent of the direction of the incident light. In the infrared, especially for $\lambda > 4 \mu$, it radiates sensibly as an ideal radiator; in the visible spectrum its emissivity is small. Its factor of entrapment (E) is about 0.05 for light, and 0.3 for total energy of sunlight; and its internal brilliance is very nearly independent of the line of sight. The attenuation is logarithmic: $T = T_0 10^{-k\pi}$, T_0 is very nearly unity, and k is about 0.1 cm⁻¹; x = depth below the surface.

Table 215.—Transmission of Radiation by Snow 375

The following data refer to direct and diffuse solar radiation; Angstrom vacuum pyranometers were used. The original paper should be consulted. τ_a = percentage of incident radiation that reached the depth d; τ_o = the corresponding percentage for the radiation that actually enters the surface. τ_o measures the true transmissivity, τ_a , the apparent transmissivity.

	Unit of $d=1$	cm; of $\tau = 1$ per cent	
Snow→		Dry —	Wet
d	τ_a	τ_c	τ_{σ}
2.5	16.0		
5	7.5		8.0
10	2.3	18.5	2.4
15	1.3	5.5	1.1
20	1.0	3.2	
25	0.8	2.2	
40	0.4	1.2	
60	0.2	0.6	

When the sky is clear, the surface temperature of snow not exposed to direct sunlight is, on account of radiation, always below that of the neighboring air, especially at high altitudes; the difference is almost as great as at night, even when the shadow is only a meter square. At night the surface temperature may be 5, 10, or even 15 °C below the temperature of the air; the lower the humidity, the greater the difference. Near midday, the surface in sunlight may melt, although the air temperature is $-10\,^{\circ}\text{C}.$ Wind reduces the difference between the temperatures of the air and the surface, and heavy cloud or fog almost obliterates it. When the sky is clear, the diurnal range in the surface temperature may amount to 20 or

^{*} Lambert's law: $i=i_0\cos\theta$, $i_0=$ intensity of radiation emitted normal to the surface, i= intensity of that emitted at an angle θ to the normal.

30 °C, even when the air temperature remains constant. The amplitude of the diurnal range in temperature decreases exponentially with the depth, and at 30 cm is of the order of 1 °C. The preceding information about snow is from J. Devaux.⁸⁵⁹ A prolonged study, extending over two years, of the temperatures at various depths in the snow blanket and in the underlying ground, at Sodankylä, Finland, within the Arctic Circle, has been made and published, with numerous citations, by J. Keränen. 856

In connection with his study of the cooling of snow during the arctic night of 1916, A. Angström 383 set up the equation H = c(dt/dx) in which H = total amount of heat, per unit surface and per unit time, received bythe snow from the air by conduction and convection, and (dt/dx) = vertical gradient of the air temperature. He called c the convectivity, and found that c = 0.005 g-cal per (°C/cm) for the average wind velocity, and that (dt/dx) = 1/12 °C/cm when the sky was clear.

Glaciers and Névés.

The emissivity of clean glaciers and névés is the same as that of ice. A névé and a surface of old, large-grained snow, are each an almost perfectly diffusing surface when clean. For a clean névé the factor of entrapment is about 0.4 for light, and 0.5 for the total energy of sunlight; for a clean glacier the corresponding values are about 0.4 and 0.6. The internal brilliance of a glacier is notably greater if the line of sight is toward the surface than for the contrary direction, but the internal illumination (I_i) is almost independent of the aspect of the surface. For a glacier, as for snow, the attenuation is logarithmic: $T = T_0 10^{-kx}$; k is a little greater for red than for green, and its value varies with the structure of the glacier, recorded values varying from 0.008 to 0.032 cm⁻¹. T₀ varies greatly with the nature of the surface, ranging from 0.2 to 0.8; it measures the attenuation produced by the surface layer. 359

J. Vallot ^{388a} has found that in the névé of the Mt. Blanc glaciers, elevation about 4.3 km, the diurnal variation in temperature does not extend below one meter, nor the annual below 6.5 meters.

EMISSIVITY OF ICE AND OF SNOW

For radiation greater than 1μ in wave-length, the absorptivity of ice, like that of water, is great (Section 75), and the reflectivity is small (Section 72); whence one may conclude that ice and snow will radiate nearly as an ideal (black body) radiator, that their emissivities will not be much less than unity (see A. Angström 888). The very low value published by K. Siegl 889 is surely incorrect.

E. Schmidt ³⁹⁰ has reported the following values for the emissivity (ϵ) of ice in terms of that of the ideal (black body) radiator, taken as unity:

Sana Vallot, J., Compt. rend., 156, 1575-1578 (1913).
 Siegl, K., Sitz.-b. Akad. Wiss., Wien (Abt. IIa), 116, 1203-1230 (1907).

son Schmidt, E., Forsch. Gebiete Ingenieurw., 5, 1-5 (1934).

Wet ice at 0 °C, $\epsilon = 0.966 \pm 0.003$; transparent ice at -9.6 °C frozen to brass, $\epsilon = 0.965 \pm 0.003$, being the same whether the thickness of the ice was 0.4 mm or 0.8 mm; white frost at -9.6 °C and 0.1 to 0.2 mm thick, $\epsilon = 0.985 \pm 0.03$, being the same whether the frost was deposited on brass or on ice.

77. Photoelectric Emission by Ice

When the illumination is that produced by the radiation from an electric spark between aluminum terminals, and filtered by a thin plate of fluorite and not more than 3 mm of air at atmospheric pressure, the photo-electric emission of electrons by ice is 280 times as great as that by water, and 0.70 times that by CuO.³⁹¹ Its variation with the filtration is shown in Table 216.

Table 216.—Relative Photoelectric Sensitivity of Ice

Adapted from A. L. Hughes ³⁹² based on the observations of W. Obolensky. ³⁹¹

Filtered radiation from an Al-spark; sensitivity = S. For the fluorite filtered radiation, the S of CuO is taken as 143. λ_{min} = shortest wavelength contained in the filtered beam.

Unit of \(\lambda\)	$= 1 \text{ m}\mu = 10\text{A} = 10^{-7} \text{ cm}$	
Filter	λm tn	S
Fluorite (CaF ₂)	125	100
Quartz (SiO ₂)	145	40
Quartz and air	177	50
Calcite (CaCO ₃)	220	0.02
Glass	330	0

[&]quot;With not more than 3 mm of air at a pressure of 1 atm.

78. Absorption Spectrum of Ice

In the region $\lambda = 6000 \text{A}$ to $6 \,\mu$, the absorption spectrum of ice is analogous to that of water, but the several wave-lengths at which the absorption passes through a maximum are each somewhat greater than the corresponding one for the liquid.³⁹³

When the optical structure of the specimen is uniform and the path of the radiation is perpendicular to the optic axis, the absorption has a maximum at each of the following wave-lengths (see E. K. Plyler 384):

Ordinary * ray,
$$\lambda = 0.79$$
 0.89 1.02 1.26 μ
Extraordinary * ray, $\lambda = 0.81$ 0.92 1.06 1.29 μ

^{*} Certain inconsistencies in this and Plyler's other paper ** reporting the same data have led to an interchange of the terms "ordinary" and "extraordinary" in several quotations of these data. The assignment here given is that in the author's Table I, which he has informed me is correct.

⁸⁹¹ Obolensky, W., Ann. d. Physik (4), 39, 961-975 (1912).

⁸⁹² Hughes, A. L., Int. Crit. Tables, 6, 68 (1929).

³⁰³ McLennan, J. C., Ruedy, R., and Burton, A. C., Proc. Roy. Soc. (London) (A), 120, 296-302 (1928); Bode, G., Ann. d. Physik (4), 30, 326-336 (1909).

In other cases, in which the radiation passed parallel to the optic axis, maxima were observed at $\lambda = 1.50$ (Bode 393 and Plyler), 1.95 and 4.5 μ (Bode).

Using a compound plate composed of portions of two crystals, Plyler observed maxima at $\lambda = 0.77$, 0.85, and 0.99 μ , the path of the radiation being perpendicular to the interface of the crystals.

The ultraviolet absorption by ice has been studied by E. J. Cassell, 894 who found a continuous absorption with a long wave-length limit near

The band near 3 μ has been studied by G. Bosschieter and J. Errera, ³⁹⁵ who found for ice only a single band with its maximum at $\lambda = 3.08 \mu$ and two inflections, one near 2.98 and the other near 3.17 μ . They ascribe the maximum (3.08 μ) to a tridymite structure in which O is surrounded by 4 H's, two being nearer the O than are the other two.

A band near $\lambda = 62 \,\mu$ has been reported for ice at $-10 \,^{\circ}$ C by C. H. Cartwright, 396 who thought that its origin is to be sought in the crystalline structure of the ice.

79. OPTICAL ROTATION BY ICE

When plane-polarized light is passed through ice-VI, the plane of polarization is rotated. 397

DIELECTRIC PROPERTIES OF ICE

The dielectric properties of ice to be considered here are its dielectric constant (ϵ') , its absorption index (κ) expressed in terms either of the equivalent conductivity $(k_c = \epsilon'' \omega/4\pi = n^2 \kappa \nu)$ or of the phase defect $(\phi = \tan^{-1}(\epsilon''/\epsilon'))$, and its dielectric strength. Symbols have been defined, dielectric theories discussed, and formulas derived in Section 49. For the electrical conductivity of ice see Section 81.

Since ice is crystalline, it is to be expected that its dielectric properties will vary with the direction that the applied field makes with the axes of the several crystals. No information bearing upon this subject has been found. All the observations seem to have rested on the tacit assumption that the axes of the individual crystals in the specimens studied had a completely random distribution. It seems improbable that the randomness was complete in any case. Differences between the results of the various observers may rest in part upon differences in the mean orientation of the axes of the crystals with reference to the field.

For theories of the structure of ice as related to the dielectric constant

⁸⁰⁴ Cassel, E. J., Proc. Roy. Soc. (London) (A), 153, 534-541 (1935).

Bosschieter, G., and Errera, J., Compt. rend., 205, 560-562 (1937); superseding Idem, 204, 1719-1721 (1937).
 See also Errera, J., Jour. de chim. phys., 34, 618-626 (1937).
 Cartwright, C. H., Nature, 136, 181 (L) (1935).

⁸⁹⁷ Poulter, T. C., Phys. Rev. (2), 37, 112 (A) (1931).

see R. H. Fowler,⁸⁹⁸ F. C. Frank,⁸⁹⁹ W. F. Giauque and J. W. Stout,⁴⁰⁰ M. L. Huggins,⁴⁰¹ C. P. Smyth,⁴⁰² A. Német.⁴⁰³

Dielectric Constant of Ice.

- O. Blüh 404 has discussed the accord between the various theories and the observed values of ϵ for a number of substances, the data for water, ice, and steam being considered in detail; a bibliography of 172 titles is given. W. Ziegler 405 also has reviewed the subject, giving a bibliography of 159 entries, and J. Errera 406 has given an exposition of theory and in the last paper cited a summary of his work on the dielectric polarization of solids.
- G. Oplatka 407 found that ice frozen from water that was not extremely well freed from gas and kept gas-free during the freezing contained large space charges, whereas pure gas-free ice contained none. Under suitable conditions the presence of a space charge may increase the effective dielectric constant 30-fold. He believed that none of the ice used by his predecessors in their study of its dielectric constant was gas-free.
- C. P. Smyth and C. S. Hitchcock ⁴⁰⁸ have reported that for ice frozen from a 0.0002M solution of KCl (1 KCl to 278 000 H₂O = 1 g KCl to 67 100 g water) ϵ' is greater than that of pure ice, and the ϵ' vs. t graphs show hysteresis at the lower frequencies, the ϵ' for increasing temperatures being less than that for decreasing. The specific conductivity (k) of the solution was $10^5k = 2.2$ (ohm·cm)⁻¹.
- E. J. Murphy 409 has found "no indication of an abrupt disappearance of the polarization responsible for the high dielectric constant of ice at any temperature above -139 °C." The main effect of lowering the temperature "appears to be an exponential increase of the relaxation time of the polarized condition of the dielectric."

In addition to the data given in the following tables and graphs, a few measurements at an unstated temperature have been reported in insufficient detail by H. Brommels.⁴¹⁰

The most extended series of measurements of ϵ' at various frequencies for ice at various temperatures are those by J. Errera, ⁴¹¹ C. P. Smyth and C. S. Hitchcock, ⁴⁰⁸ H. Wintsch, ⁴¹² and E. J. Murphy. ⁴⁰⁹ The last alone gives values for temperatures below $-70\,^{\circ}$ C, but his observations are dis-

cordant with those of the others in at least two particulars: (1) his values at -7.1 to -45.8 °C at low frequencies are much higher, ϵ' being over 95

Table 217.—Drude-Debye Constants for the Dielectric Constant of Ice

$$\epsilon = \epsilon' - i\epsilon'', \ \epsilon' = \epsilon_0 + (\epsilon_1 - \epsilon_0)/(1 + a^2v^2); v_s = 1/a, \lambda_s = ca, \tau = \frac{a}{2\pi} \times (\epsilon_0 + 2)/(\epsilon_1 + 2), \lambda_s = \text{transition wave-length.}$$
 See Section 49, p. 356. It has been found empirically that $a = \alpha e^{-\beta t}$. Each of the three extended series of observations now available lead to a different set of values for ϵ_0 , ϵ_1 , α , and β , the most consistent being those by SH.⁴⁰⁸

Unit of a and of $\tau = 1$ sec; of $\nu_s = 1$ cycle/sec; of $\lambda_s = 1$ km. Temp. = t °C

I. Smyth and Hitchcock (1932).⁴⁰⁸ $\epsilon_0 = 3.0$, $\epsilon_1 = 74.6$, $\alpha = 116.0$ microsec; $\beta = 0.1015$ (°C)⁻¹.

-0	1.160	1.346	8620	34.8	1.205
-2	1.421	2.019	7037	42.6	1.48
 5	1.927	3.713	5189	57.8	2.00
-10	2.202	4.849	4541	66.1	2.29
-15	5.217	27.22	1917	156.5	5.42
-20 -25 -30 -40	8.833	78.02	1126	265	9.17
-25	14.66	214.6	682	440	15.23
-30	24.37	593.9	410	731	25.3
-40	67.23	4520	149	2017	69.9
-50	185.6	34450	53.9	5560	193
-60	512.1	262200	19.52	15360	532
-70	1413	1997000	7.08	42360	1467

II. Wintsch (1932).⁴¹² $\epsilon_0 = 7.5$, $\epsilon_1 = 73.0$, $\alpha = 141.2$ microsec., $\beta = 0.0906$ (°C)⁻¹.

-0	1.412	1.994	7082	42.3	2.84
-5	2.221	4.933	4502	66.6	4.48
-10	3.492	12.194	2864	104.7	7.04
-20	8.64	74.65	1157	259	17.40
-30	21.20	449.4	472	636	42.7
-40	52.95	2803.7	188.8	1587	106.6
-50	130.3	16980	76.7	3907	262,4

III. Errera (1924).⁴⁰⁶ $\epsilon_0 = 3.0$, $\epsilon_1 = 77.2$, $\alpha = 182$ microsec., $\beta = 0.090$ (°C)⁻¹.

-0	1.82	3.312	5491	54.6	1.83
$-2 \\ -5$	2.178	4.744	4591	65.2	2.19
-5	2.855	8.151	3503	86.5	2.87
-10	4.477	20.04	2248	134.2	4.50
-20	11.01	121.2	908	330	11.07
$-22 \\ -25$	13.20	174.1	758	396	13.3
-25	17.27	298.2	579	518	17.4
30	27.08	733.4	369	812	27.2
-37	50.9 6	2597	196	1528	51.2
-40	66.61	4437	150	1998	66.9
-50	164.0	26900	61.0	4914	165

IV. Murphy $(1934)^{400}$ gives $10^5 \tau_M = 1.85e^{-0.106t}$ where $\tau_M = \tau(\epsilon_1 + 2)/(\epsilon_0 + 2) = a/2\pi$. Whence $10^6 a = 116e^{-0.106t}$. See remarks in text.

Table 218.—Dielectric Constant of Ice: Observed and Computed

(See Table 219 and Figures 10 and 11 for other observed values.)

 $\epsilon' = 3.0 + 71.6(1 + a^2 v^2)^{-1}$, v = frequency, $10^4 a = 1.160 e^{-0.1015t}$ sec. Temp. = t °C; Dif $\equiv \text{Obs.} - \text{Calc.}$ The ice was Observed (Obs) values are from C. P. Smyth and C. S. Hitchcock 408; computed (Calc) values are those defined by formed of "conductivity water" for which $10^6 k = 2 (\text{ohm cm})^{-1}$.

2.99 3.00 -0.01 3.33 3.00 +0.33 2.97 3.00 -0.03 3.59 3.01 +0.58 $3.12 \quad 3.00 \\ +0.12$ 3.00 3.00 3.50 3.03 +0.47 $3.92 \quad 3.11 \\ +0.81$ 3.00 3.00 262246 3.04 3.00 +0.04 $3.02 \quad 3.00 \\ +0.02$ $6.17 \quad 5.24 \\ +0.93$ 4.65 3.82 +0.83 Obs Cale $3.21 \quad 3.01 \\ +0.20$ 2 3.21 +0.61 34447 3.82 Unit of a=1 sec; of $\nu=1$ kilocycle/sec; of $\epsilon'=1$ cgse unit. Temp. = t °C 8.82 4.55 3.06 3.00 3.03 3.00 Obs Calc Dif 16.3 17.1 +0.82 4520 5.37 3.32 8.78 3.48 3.03 $31.3 \quad 31.8 \\ -0.5$ 3.04 3.00 $14.6 \quad 13.3 + 1.3$ Obs Calc Dif 49.7 593.9 -3046.2 4 -3.5 45.2 43.2 +2.0 7.60 6.49 + 1.11Obs Calc 65.9 62.9 71.9 78.022 5.46 & 6.51 -1.053.33 3.41 -0.0873.6 73.8 69.4 71.2 74.8 74.3 +0.5 24.4° 35.3 -10.9 Obs Cale Dif 4.8488 · Though so printed, this value seems to be 8.34 7.52 +0.82 $\frac{3.71}{+0.18}$ 73.8 74.0 -0.2 72.5 72.0 +0.5 2 40.1 +0.1 Obs Calc Dif 3.7133 5 40.2 4.2 3.78 +0.4 10.4 9.6 +0.8 74.2 74.4 -0.2 73.5 74.2 -0.7 72.4 72.9 -0.5 46.3 47.2 Obs Calc Dif 2.4743 4.3 4.18 + 0.112.3 12.4 73.6 74.3 -0.7 72.5 73.4 -0.9 51.6 53.7 1.6512 $10^8 a^2 \rightarrow$ 0.5 0.3 ន 8

Table 219.—Dielectric Constant of Ice

(See also Figures 10 and 11.)

At -5 °C $\epsilon' = -0.08 + 0.34 \log_{10^{\nu}}$ cgse units, if $10^7 < \nu < 10^9$ cycles/sec. ⁴¹³ E. J. Murphy ⁴⁰⁹ has reported the following high values, as read from his graphs, unit of ν being 1 cycle/sec: -90 °C, ballistic method, $\epsilon' = 150$; -45.6 °C, $\nu = 15$, $\epsilon' = 87$; -7.1 °C, $\nu = 300$, $\epsilon' = 95$; for other values, consult his paper.

Unit of $\lambda = 1$ km; of $\nu = 1$ kilocycle/sec; of $\epsilon' = 1$ cgse. Temp. = t °C

- I. Smyth and Hitchcock, 1932. See Table 218.
- II. Wintsch,^a 1932, read from his graphs. Water was thrice distilled, collected in quartz, and boiled just before freezing.

$t \rightarrow \nu$	-5	-6	-10	-20	-30	-40	-50	
0.05	85.3		85.2	82.0	74.0			
0.65	74.0		71.2	58.5	37.0	21.8	17.0	
1.00		69.7	65.5	44.5	23.5	15.1	12.4	
1.13	69.2		64.5	41.8	21.8	14.2	11.2	
1.60	65.5		57.0	30.0	16.0	11.0	9.3	
2.00		59.5	51.4	23.6	12.9	9.6	8.5	
3.00		48.9	38.7	16.4	10.0	8.0		
3.50	46.5		33.3	13.8	8.7	7.5	7.5	
4.00		40.2	29.6	12.4	8.9	7.3		
5.00		33.2	24.0	10.2	8.0	6.8		
5.10	34.8		23.0	10.0	7.0	6.3	6.2	
6.00		28.0	19.4	9.0	7.1			
7.00		24.1	16.4					
8.00		21.0	14.5					

III. Errera, a 1924, certain typographical errors corrected.

$\lambda \rightarrow \lambda$		-2	-5	-226	-370	-47.5
680	0.441	77.3	76	43.5		7.4
465	0.645	76	74	34.3	10.15	4.15
430	0.698		• •	31.5	8.3	3.68
294	1.020	73.4	72.6	19.4	4.6	3.22
196	1.53	69	65.8	13	4.15	
79ª	3.80	47.2	39.8	5.06		
54	5.56	30.6	25.4	4.6	3.2	2.76
38.5	7.80	23.2	16.6	3.4	3	
28.5	10.5	15.2	11.6	3.1		
18.5	16.2	7.8	5.6			
8	37.5	4.6	4.4	2.3		2.3
1.1	273		3.86	2.2		

IV. Granier,^a 1924, t = -12 °C: from water having $10^6 k = 1.54$ (ohm·cm)⁻¹.

λ	ν	€′	λ	ν	e'
70000	0.0043	153	17.6	17	3.8
6000	0.050	100	5.9	51	2.35
940	0.320	86	1.15	260	2.05
194	1.55	56	0.045	6700	2.05
56	5.40	12			

618 Curtis, H. L., and Defandorf, F. M., Int. Crit. Tables, 6, 78 (1929); from Gutton, C., Compt. rend., 130, 1119-1121 (1900).

Table 219—(Continued)

V. Adapted from International Critical Tables; CD.a

$t\rightarrow$	ϵ'	*	ϵ'	t ε'
$\nu = 0.050$;	Ths ^a	-70	41.5	$\nu = 0.320$; DF'
-2	94	-80	31.5	-7 51
-10	95.2	-90	20.2	-47 3.6
-18	96.5	-100	14.5	$\nu = \text{audio}; A^a$
-182	3.	-110	8.6	-80 3.8'
$\nu = 0.120^{\circ}$	DF, FD ^a	-120	6.1	$\nu = 10000$; Thg ^a
-20	59.5	-130	4.7	-2 3.4
-30	59.0	-140	3.5	-5 2.8°
-40	58.5	-150	2.7	$\nu = 100000$; BK ^a
-50	56.0	-165	2.43	-190 1.8 ^a
-60	49.5	-185	2.43	

* References:

- A BK
- Abegg, R., Ann. d. Physik (Wied.), 62, 249-258 (1897).
 Behn, U., and Kiebitz, F., "Boltzmann Festschrift," p. 610-617, 1904.
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 Dewar, J., and Fleming, J. A., Proc. Roy. Soc. (London), 61, 2-18 (1897).
 Dewar, J., and Fleming, J. A., Idem, 62, 250-266 (1898).
 Errera, J. 411 CD
- DF'
- Er Eleming, J. A., and Dewar, J., *Proc. Roy. Soc. (London)*, **61**, 316-330 (1897). Granier, J., *Compt. rend.*, **179**, 1313-1318 (1924). Thwing, C. B., *Z. Physik. Chem.*, **14**, 286-300 (1894). Thomas, P., *Phys. Rev.*, **31**, 278-290 (1910). Wintsch, H.⁴¹² FD

- Ths

^b All of these -22 °C values are out of line with the others (see Fig. 9); it seems probable that the temperature is misprinted. If it was actually somewhere between -25 and -27 °C the values would about fit.

This temperature was printed -27 °C, but -37 °C makes these values fit with

the others; probably a misprint. "This λ was printed as 97, but 79 is required to bring the values into line with the others. The ν 's have been computed by the compiler, and the 3.80 corresponds to the 79.

The condenser was charged and discharged 120 times a second by means of a

vibrating contact-maker controlled by a tuning fork.

'At 5 megacycles/sec this same value (3.8) was found throughout the range 0 to -24 °C.⁴¹⁴

'At -5 °C and 10 megacycles/sec C. Gutton ⁴¹³ found $\epsilon'=2.3$.

At -4.5 °C and 83 megacycles/sec B. de Lenaizon and J. Granier ⁴¹⁵ found $\epsilon'=1.5$

2.17.

at -7.1 °C and nearly 90 at -45.8 °C, and his ballistic values at -90 °C are about 150; (2) his graph shows that the logarithm of Debye's 1 (see page 355 is linear in the reciprocal of the absolute temperature, whereas the other sets of data just mentioned indicate that it is linear in the temperature. True, Murphy gives for τ an expression that requires the logarithm to be linear in the temperature, but that expression is a mere approximation, compromising with the graph and limited by him to temperatures above -46 °C. It should be noticed that his τ is so defined as to be $(\epsilon_1 + 2)/(\epsilon_0 + 2)$ times as great as Debye's.

From the data by each of the other three, the present compiler has determined graphically and by cut-and-try methods the three constants $(\epsilon_0, \epsilon_1 - \epsilon_0, \text{ and } a)$ occurring in the Drude-Debye isothermal relation $\epsilon' =$ $\epsilon_0 + (\epsilon_1 - \epsilon_0)/(1 + a^2\nu^2)$ (see eq. 9, Section 49), and has found in each

⁴¹⁴ Abegg, R., Ann. d. Physik (Wied.), 65, 229-236 (1898).

⁴¹⁸ de Lenaizon, B., and Granier, J., Compt. rend., 180, 198-199 (1925).

case that the logarithm of a is linear in t; a is proportional to Debye's τ . The values of all three sets of constants and of the α and β in $\log a = \log \alpha - \beta t$ are given in Table 217. It will be noticed that the three β 's

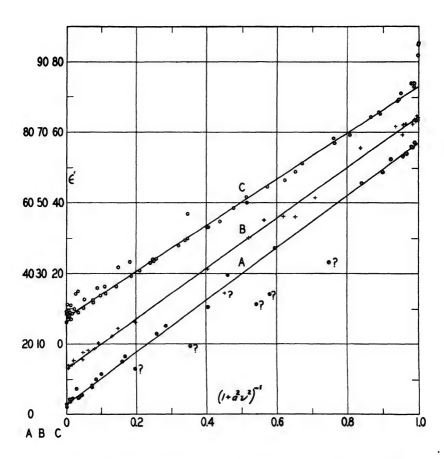


Figure 9. Dielectric Constant of Ice: Variation of ϵ' with $(1 + a^2 r^2)^{-1}$.

The observed values of ϵ' given by Errera (A), Smyth and Hitchcock (B), and Wintsch (C), and contained in Tables 218 and 219 are plotted against the reciprocal of $(1 + a^2p^2)$, the value of a being in each case determined from the constants given in Table 217. The origin of the scale of ϵ' is shifted from curve to curve, each scale being appropriately marked. The 5 questioned values of Errera's all refer to the temperature published as -22 °C; if the actual temperature was somewhere between -25 and -27 °C these points would lie near the line.

differ but little, but the α 's differ greatly, suggesting that the three samples of ice differed significantly in some manner. It seems possible that some

single value of β might be used in all three cases, and the α 's be adjusted so as to obtain a satisfactory agreement with the observations, but time for testing this is not now available. It will also be noticed that, as in the case of water, the values that must be used for ϵ_0 and ϵ_1 do not agree respec-

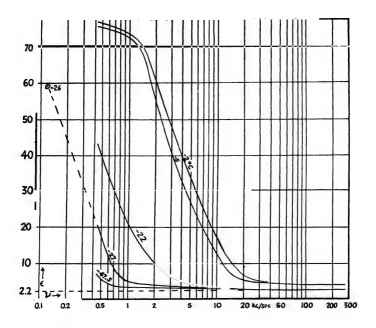


FIGURE 10. Isothermal Variation of the Dielectric Constant (ϵ') of Ice with the Frequency (ν) of the Field.

Unit of $\epsilon' = 1$ cgse; of $\nu = 1$ kilocycle/sec.

[Adapted from compilation by H. L. Curtis and F. M. Defandorf, Int. Crit. Tables, 6, 78 (1929)—based upon data by J. Errera, Jour. de Phys. (6), 5, 304-311 (1924)—with the addition of an observation (circle) by J. A. Fleming and J. Dewar, Proc. Roy. Soc. London, 61, 316-330 (1897).]

tively with the square of the optical index and with the static value of ϵ' . After the values of the constants had been obtained, each value of ϵ' of each set was plotted against the reciprocal of the corresponding value of $(1 + \alpha^2 v^2)$, using for α the value defined by the derived values of α and β . These graphs are shown in Figure 9. It will be noticed that all the values of ϵ' given by Smyth and Hitchcock, whatever the temperature and fre-

quency may be, lie quite satisfactorily along a right line, with a single exception. Those read from Wintsch's graphs do likewise except at the extremities, but the spread is greater; and Errera's values, corrected for two obvious misprints (see notes to Table 219) fit well, excepting the set

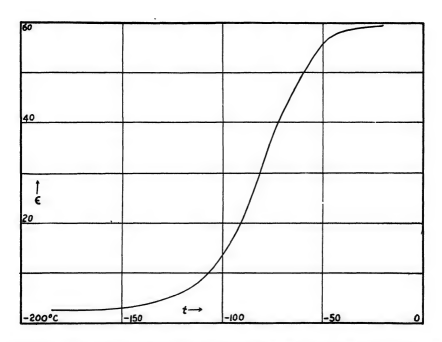


FIGURE 11. Thermal Variation of the Dielectric Constant (ε') of Ice for 120 Charges and Discharges per Second.

[From compilation by H. L. Curtis and F. M. Defandorf, Int. Crit. Tables, 6, 79 (1929), based upon data by J. Dewar and J. A. Fleming, Proc. Roy. Soc. London, 61, 2-18 (1897), and by J. A. Fleming and J. Dewar, Idem, 61, 316-330 (1937). Cf. Table 219, Section V.]

for $-22\,^{\circ}$ C, which are consistently lower. If the temperature given in his table as $-22\,^{\circ}$ C was actually somewhere between $-25\,$ and $-27\,^{\circ}$ C, these points also would lie close to the line. It seems probable that this is another misprint. In Table 218 each of the values of ϵ' given by Smyth and Hitchcock is compared with the corresponding one as computed from the constants obtained from their complete set of values.

Dielectric Absorption of Ice.

(For definitions of terms and symbols see Section 49.) The value of the dielectric absorption of ice is commonly indicated by means of either the phase defect $\phi = \tan^{-1}(\epsilon''/\epsilon')$ or the apparent conductivity $k_a = \epsilon'' \nu/2$ cgse units = $(c^2 \epsilon'' \nu/2) 10^{-9} (\text{ohm-cm})^{-1}$, ϵ'' being expressed in cgse units, and c = velocity of light.

The values of ϵ'' and of ϵ' can be computed from the constants given in Table 217, and from them ϕ and k_a may be obtained. These calculations have been carried through for the observations of Smyth and Hitchcock and are given in Tables 220 and 222. Whether the experimentally determined values of ϕ and k_a are entirely free from the effects of such true conductivity as the ice may have had, is not entirely clear.

Table 220.—Phase Defect for Ice: Observed and Computed

 $\phi=\tan^{-1}(\epsilon''/\epsilon')$. Calculated (Calc) values are those defined by the formulas $\epsilon'=\epsilon_0+(\epsilon_1-\epsilon_0)/(1+a^2\nu^2)$ and $\epsilon''=(\epsilon_1-\epsilon_0)a\nu/(1+a^2\nu^2)$ with $\epsilon_0=3.0$, $\epsilon_1=74.6$; $10^4a=1.160e^{-0.1015t}$ sec. Temp. =t °C. See text. Observed (Obs) values are those derived from the values given by C. P. Smyth and C. S. Hitchcock ⁴⁰⁸ for the equivalent conductivity and the dielectric constant. The specific electrical conductivity (k) of the water used was $10^6k=2(\text{ohm}\cdot\text{cm})^{-1}$.

Unit of y = 1 cycle/sec: of $\phi = 1^\circ$ of arc. Temp = $t^\circ C$

		Unit of $\nu = 1$ cyc	cle/sec; of $\varphi = 1$	of arc. 1cm	$p_{i} = t^{\circ}C$	
$\nu/1000 \rightarrow$	0.3	0.5	1 · · · · · · · · · · ·	5	20	60
t	Obs Calc Dif	Obs Calc Dif	Obs Calc Dif	Obs Calc Dif	Obs Calc Dif	Obs Calc Dif
-1	4.0 2.3 + 1.7	$4.8 3.5 \\ +1.3$	8.0 7.0 +1.0	$31.2 31.6 \\ -0.4$	61.8 62.8 - 1.0	64.4 65.5 -1.1
-3	$\begin{array}{ccc} 2.4 & 2.6 \\ -0.2 & \end{array}$	$\begin{array}{c} 4.8 & 4.3 \\ +0.5 \end{array}$	$9.2 8.6 \\ +0.6$	$36.8 36.3 \\ +0.5$	$63.7 65.0 \\ -1.3$	$63.7 63.3 \\ +0.4$
-5	$3.3 3.2 \\ +0.1$	5.6 5.3 + 0.3	$11.4 10.5 \\ +0.9$	$41.4 41.7 \\ -0.3$	64.8 66.6 -1.8	$62.4 60.1 \\ +2.3$
-10	$\frac{4.6}{+1.0}$ 3.6	$11.1 6.0 \\ +5.1$	$18.0 11.9 \\ +6.1$	53.0 45.2 +7.8	$64.4 ext{ } 67.2 ext{ } -2.8$	54.5 57.6 -3.1
-20	$14.5 14.2 \\ +0.3$	25.2 22.8 +2.4	$40.4 39.4 \\ +1.0$	65.1 67.1 -2.0	$54.2 51.4 \\ +2.8$	$32.8 24.0 \\ +8.8$
-30	38.9 34.5 +4.4	53.6 47.8 +5.8	$61.2 62.1 \\ -0.9$	58.5 59.2 •−0.7	$30.3 25.9 \\ +6.4$	18.2 9.3 +8.9
-40	60.5 59.0 +1.5	$66.1 65.8 \\ +0.3$	$62.7 66.4 \\ -3.7$	$36.8 34.7 \\ +2.1$	14.7 10.0 +4.7	$14.3 3.4 \\ +10.9$
-50	63.9 67.1 -3.2	58.4 63.4 -5.0	48.5 50.2 -1.7	18.0 14.4 +3.6	$7.3 3.7 \\ +3.6$	$7.7 ext{ } 1.2 ext{ } +6.5$
-60 ·	49.8 54.6 -4.8	38.3 41.9 -3.6	29.5 24.8 +4.7	10.4 5.3 + 5.1	$^{4.1}_{+2.8}$	6.8 0.4 +6.4
-70	38.0 29.1 +8.9	27.5 18.6 +8.9	20.4 9.6 +10.8	7.6 1.9 +5.7	$1.9 0.5 \\ +1.4$	6.4 0.2 +6.2

Table 221.—Phase Defect for Ice

For definitions of terms and symbols see Section 49; $\tan \phi = \epsilon''/\epsilon'$. The electrical conductivity (k) of the water used by Wintsch was not stated; it was triply distilled and collected in quartz. For that used by Granier $10^6 k = 1.5 (\text{ohm} \cdot \text{cm})^{-1}$.

Unit of v = 1 kilocycle/sec; of $\phi = 1^{\circ}$ of arc. Temp. = t° C

I. Smyth and Hitchcock, 1932, see Table 220.

II.	Wintsch,412	read from	his	graphs.
-----	-------------	-----------	-----	---------

$t \rightarrow$	-5	-6	-10	-20	-30	-40	-50	
0.05	1.2	40.4	1.7	3.4	8.5	40.0		
0.5 0.650	12.2	10.4	13.0 15.0	24.5 29.0	40.0 43.5	48.0 51.3	51.7 53.5	
1.0	10.0	16.4	20.8	37.7	51.0	54.3	54.3	
1.13	16.4		22.0	40.0	52.2	54.0	53.0	
1.60	20.2		28.5	52.5	55.1	53.0	51.0	
2.0		26.0	33.1	51.5	55.8	52.5	49.0	
3.0 3.5	36.0	34.6	42.0 46.0	56.4 57.5	53.3 53.5	47.8 45.8	43.2 40.5	
4.0	00.0	41.0	47.8	58.1	52.7	44.4	39.8	
5.0		45.7	52.1	59.0	51.2	41.5	38.0	
5.1	44.5	10.4	52.7	59.1	51.0	41.0	37.5	
6.0		49.6	55.6	59.5				
7.0		52.5	58.0					
8.0		54.0	58.8					
III. G	iranier,410	1924; t	= -12	°C.				

ν	0.050	0.320	1.550	5.40	17.0	51.0	260	6700	64000
		32							

^a This value (0.5) is from Granier.⁴¹⁷

Dielectric Strength of Ice.

P. Thomas 417a has found that in a uniform alternating field applied at the rate of about 600 volts per second (frequency = 1000 cycles/sec) ice broke down when the field reached the value of 11,000 volts/cm; $\epsilon' = 86.4$ cgse, conductivity = $1.4 \times 10^{-8} (\text{ohm} \cdot \text{cm})^{-1}$; temperature is not stated.

81. ELECTRICAL CONDUCTIVITY OF ICE

In any discussion of the electrical conductivity of ice it is quite essential to recognize the several distinct types of effect that contribute to the observed effective conductivity.

In most cases, the value assigned to the (effective) conductivity is that derived from the resistance R which must be placed in parallel with a pure capacitance C in order to obtain an exact equivalent of the actual icecondenser under the existing conditions. If the ice-condenser were merely a leaky condenser—if the ice were composed of an ideal, nonconducting,

⁴¹⁶ Granier, J., Compt. rend., 179, 1313-1318 (1924).

⁴¹⁷ Granier, J., Bull. Soc. Fr. des Elec. (4), 3, 333-482 (1923).

⁴¹⁷a Thomas, P., J. Franklin Inst., 176, 283-301 (1913).

Table 222.—Apparent Electric Conductivity of Ice: Observed and Computed

defined by the formulas $\epsilon'' = (\epsilon_1 - \epsilon_0)a\nu/(1 + a^2\nu^2) = (\epsilon_1 - \epsilon/)/a\nu$ and $k_a = \nu\epsilon''/2$ cgse $= (\nu\epsilon''/18) \times 10^{-11} (\text{ohm-cm})^{-1}$ in which $\epsilon_0 = 3.0$, $\epsilon_1 = 74.6$, $10^4a = 1.160e^{-0.1016t}$ sec. Temp. = t °C, and $\epsilon' = \epsilon_0 + (\epsilon_1 - \epsilon_0)/(1 + a^2\nu^2)$. See text of Section 49. The electrical conductivity (k) of the water was $10^6k = 2(\text{ohm-cm})^{-1}$. Observed (Obs) values are those reported by C. P. Smyth and C. S. Hitchcock 408; calculated ones (Calc) are those

			Unit	Unit of $k_a = 1$ (ohm.cm) ⁻¹ ; of $\nu = 1$ cycle/sec.	$^{-1}$; of $\nu = 1$	cycle/sec. Ten	Temp. = t °C			
1	-1	-3	<u>ن</u> ا	-10	-20 108b.	-30	-40	-20	09-	-70
/1000	/1000 Obs Calc	Obs Calc Dif	Obs Calc Obs Calc O	Obs Calc Obs Calc O	Obs Calc Dif	Obs Calc Dif	Obs Calc Dif	Obs Calc Dif	ic Obs Calc Obs Calc Obs Calc Obs Calc Obs Calc Dif Dif Dif Dif	Obs Calc Dif
0.3	0.086 0.046	0.052 0.056	0.07 0.069 0.00	0.10 0.078 + 0.02	$0.31 0.296 \\ +0.01$	$0.62 0.570 \\ -0.05$	0.48 0.474 +0.01	$0.21 \begin{array}{c} 0.207 \\ 0.0 \end{array}$	0.086 0.077 +0.009	0.052 0.028 +0.024
0.5	$0.17 0.127 \\ +0.04$	$0.17 \begin{array}{c} 0.155 \\ +0.01 \end{array}$	$0.20 & 0.190 \\ +0.01$	$0.40 0.216 \\ +0.18$	$0.86 0.735 \\ +0.12$	$1.18 0.974 \\ +0.21$	0.55 0.544 +0.01	$0.21 0.212 \\ 0.0$	0.086 0.078 +0.008	0.052 0.028 +0.024
-	0.57 0.503	$0.65 0.611 \\ +0.04$	$0.81 0.739 \\ +0.07$	1.25 0.834 +0.42	2.14 1.97 +0.17	$1.48 1.40 \\ +0.08$	0.58 0.579 0.0	$0.24 0.214 \\ +0.03$	$0.11 0.078 \\ +0.03$	$0.069 0.028 \\ +0.041$
NO.	8.85 9.05 -0.20	9.60 9.65	9.86 9.93 -0.07	9.0 9.88 -0.88	4.55 4.28 +0.27	$1.84 1.62$ ± 0.22	$0.69 0.59 \\ +0.10$	0.29 0.214 +0.08	$0.16 & 0.078 \\ +0.08$	$0.11 0.028 \\ +0.08$
20	$\begin{array}{ccc} 25.6 & 26.9 \\ -1.3 & \end{array}$	23.3 22.9 +0.4	$19.7 19.4 \\ +0.3$	12.7 17.2 -4.5	5.5 4.49	2.06 1.63 +0.43	0.89 0.59 +0.30	$0.43 0.214 \\ +0.22$	$0.24 & 0.078 \\ +0.16$	$0.11 0.028 \\ +0.08$
9 '	29.9 30.5 -0.6	27.5 25.0 +2.5	23.7 20.5 +3.2	15.6 18.0 + 7.6	6.7 4.50 +2.2	$3.33 1.63 \\ +1.70$	2.58 0.59 +1.99	$1.37 0.214 \\ +1.16$	$^{1.2}_{+1.1}$	$^{1.1}_{+1.1}$ 0.028

For additional data and remarks see Section 81.

electrically perfectly elastic dielectric interpenetrated along the lines of electric force by threads of an ideal conductor—then R would be the combined resistance of those threads, and the conductivity computed from it would be the actual conductivity of the ice, which may be designated by k.

But the problem is not so simple. Unless the ice is exceedingly pure and gas-free, which last has probably not been the case in any of the work, it will acquire a space-charge under the action of the field. With constant fields this will act as a kind of polarization, adding itself to the polarization of the electrodes. With alternating fields, it may constitute a quite significant part of the actual current. The energy dissipated by the alternating concentration of this charge now nearer one electrode and now nearer the other will contribute to R, giving rise to a new term (k_o) in the effective conductivity. It seems reasonable to expect that both k and k_o will steadily decrease as the temperature falls, and that for a given temperature, k_o will increase with the frequency (ν) , but k will not.

Furthermore, the molecules of H_2O are polar. Hence, they will tend to place their electrical axes parallel to the impressed field.* The dissipation involved in such reorientation will also contribute to R, and hence to the effective conductivity. $^{418, \text{ pp. }80\text{-}108}$ Denote this component by k_p . As the temperature is decreased, it is to be expected that both the resistance to the rotation involved in such reorienation will be increased and the extent of the rotation will be decreased. The former will increase, and the latter will decrease, the dissipation. Whence, one should expect the dissipation, and consequently k_p , to pass, in general, through a maximum, and then to decrease to zero, the frequency being constant and not too great. If the frequency is very high, the molecules may not have time to rotate through an appreciable angle, and k_p will be zero. (See also p. 504.)

Thus it is evident that the effective conductivity (k_e) of ice is, in general, made up of at least three terms, $k_e = k + k_c + k_p$. The static conductivity is k, suitable correction being applied for such polarization as may exist; at intermediate frequencies, k_p may be the dominant term; and at high frequencies, k_c may be supreme. The data at present available do not suffice for a complete separation of these three terms, but they are consistent with the ideas just expressed (see Table 223). For example, Johnstone's static values for ice $(10^8k = 2.80 \text{ (ohm·cm)}^{-1} \text{ at } 0 \text{ °C}$ and 0.026 at -19 °C, electrolytic polarization eliminated) are not only much lower than the conductivity $(10^8k = 71 \text{ at } 17.9 \text{ °C})$ of the water from which the ice was frozen, but are also lower than that $(10^8k = 4)$ of the purest water obtained by Kohlrausch. On the other hand, Smyth and Hitchcock found at 60 kc/sec and -1 °C the great value $10^8k_e = 29.9$, over 7 times that of the purest water, and actually 15 per cent of that $(10^8k = 200)$ of the water from which it was frozen; the value decreased continuously with the tem-

^{*}P. Debye 418 has shown that even if the maximum dielectric constant (80) were due entirely to such orientation only 1 molecule in 5 million need follow the field.
418 Debye, P., "Polar Molecules," p. 106, New York, Reinhold Publishing Corp., 1929.

perature. It seems that here k_{σ} is the dominant term. In the range $\nu = 0.3$ to 1 kc/sec, the same authors found that k_{σ} passes through a maximum, the value at the maximum decreasing with ν . Thus, at $\nu = 1$ kc/sec, $10^8k_{\sigma} = 0.57$ at -1 °C, 2.14 (max.) at -20 °C, and 0.069 at -70 °C, all lower than that for the purest water. Here k_p is dominant. Why the values at -1 °C and $\nu = 0.3$ to 1 kc/sec are so much lower than the static one found by Johnstone at 0 °C is not clear.

For a complete interpretation of the variation of k_{θ} with the tempera-

Table 223.—Electrical Conductivity of Ice (See also Table 224.)

The text should be consulted. The conductivity of the water from which the ice was formed is indicated in each case. It seems probable that the k_0 of SH and of G is essentially k_0 .

	Unit of	k = 10-8	(ohm cm)	-1; of v =	= 1 kilo	cycle/sec	. Tem	$p. = t^{\circ}$	<u> </u>	
		Stati	c: kwate	r = 71° at	17.9 °	C (ICT	() 0			
	t	0 2.8		$-4 \\ 0.23$	•	-10 0.11		-19 0.0	26	
ν →	0.3	3	0.5	1		5		20	6	0
t				kwate	r = 200 ke	(SH)				
-1 -3 -5 -10	0.08 0.05 0.07	52 7	0.17 0.17 0.20	0.57 0.65 0.81		8.85 9.62 9.86	1	25.6 23.3 9.7	29. 27. 23.	.5 7
$-10 \\ -20$	0.10		0.40 0.86	1.25 2.14		9.0 4.55		2.4 5.5	15. 6.	
-30 -40 -50	0.62 0.48 0.2	3	1.18 0.55 0.21	1.48 0.58 0.24		1.84 0.69 0.29		2 06 0.89 0.43	2.	.33 .58 .37
-60 -70	0.08		0.086 0 052	0.11 0.06	9	0.16 0.11		0.24 0.11	1	2 .1
t = - 12 °C				kwate	r = 154	(G) b				
k. °	0 0.004	0.0043 0.02	0.050 0 13	0.320 1.09	1.55 6 0	5.40 9.1	17.0 9.4	51.0 9.4	260 9.8	6700 9.4

^a See J. H. L. Johnstone. ⁴¹⁹

ture, consideration must be given to the progressive melting discussed by Buchanan (p. 469).

J. H. L. Johnstone 419 has stated that the effects of polarization are great, are not entirely electrolytic, and are difficult to eliminate. The observations of G. Oplatka 407 indicate much the same. Johnstone used potential leads, and measured their potentials electrostatically. The only measurements at exceedingly low temperatures seem to be those by Dewar

^{*} References:

G Granier, J.⁴¹⁶
ICT Bjerrum, N.⁴²² Based on observations of J. H. L. Johnstone.⁴¹⁶
SH Smyth, C. P., and Hitchcock, C. S.⁴⁰⁸

^o Computed by the compiler from Granier's data.

⁴¹⁹ Johnstone, J. H. L., Proc. Trans. Nova Scotian Inst. Sci., 13, 126-144 (1912).

⁴²⁰ Fleming, J. A., and Dewar, J., Proc. Roy. Soc. (London), 61, 316-330 (1897).

and Fleming (Table 224). The method employed is not described, but it appears to have been a static one, the resistance being derived by means of Ohm's law from the steady current and the applied voltage. There is no

Table 224.—Thermal Variation of the Electrical Resistance of Ice

The several sets of values of R are not comparable.

It seems probable that the values attributed to Wintsch (read from his graphs) represent mainly the component (k_p) arising from dielectric absorption. Those attributed to Dewar and Fleming seem to have been inferred from the constant impressed voltage and the observed current; the values are merely approximate, and the two sets, having been obtained with different vessels, are not comparable; the first set refers to ice from ordinary distilled water, the second to that from especially pure water; t_p is the temperature inferred from their platinum thermometer, and is lower than the actual temperature on the centigrade scale. Frequency is ν .

Unit of R=1 ohm; of $\nu=1$ kilocycle/sec. Temp. = t °C

l. H.	Wintsch.	412					
$t \rightarrow$	-5	-6	-10	-20	-30	-40	-50
ν				R/1000			
0.05	57500		42400	19000	7750		
0.65			342	202	184	233	278
1.00		198	172	118	139	202	249
1.13	184		144	108	133	194	240
1.60	112		86	81	116	180	224
2.00		76	65	70	111	172	216
3.00		42	42	59	102	159	198
3.50	36		35	56	99	152	192
4 00		31	32	53	96	148	182
5.00		24	28	49	92	139	168
5.10	23		28	49	92	138	178
6.00		20	26	48			•

II. J. Dewar and J. A. Fleming. 423 Fleming and Dewar. 420

-	Dis	tilled			Pu	re	
t_p	R/106	t_p	R/10 ⁸	t_p	R/106	t_{P}	R/10°
-70.7	43.4	-93.2	282	-10.2	1	-92.0	1200
-75.0	42.8	-95.2	353	-19.2	3	-138.2	2000
-82.3	46.3	-98.8	470	-26.1	15	-152.1	2500
-84.4	53.4	-108.4	706	-27.6	40	-206	25000
-86.3	66.5	-126.0	1130	-33.2	250		
-88.2	91.4	-135.0	1570	-42.1	260		
-88.8	118.0	-172.0	5670	-47.0	410		
-91.9	209	-200.0	26200	-68.2	1200		

way in which the actual conductivity can be inferred from the resistances they tabulated. They remark: "Above a certain temperature there is a relatively rapid increase in the conductivity of the ice, as it rises in temperature." 420 A single series of observations at an unstated frequency has

⁴²¹ Brommels, H., Comment. Phys.-Math. Soc. Sci. Fennica (Helsingfors), 1, No. 19 (1922).

⁴²² Bjerrum, N., Int. Crit. Tables, 6, 152 (1929).

⁴²⁸ Dewar, J., and Fleming, J. A., Proc. Roy. Soc. (London), 61, 2-18 (1897).

been published in insufficient detail by H. Brommels, 421 the values of 10^8k_e increasing from 38.9 at $-1.2\,^{\circ}\text{C}$ to 47.3 at $-17.3\,^{\circ}\text{C}$. H. Wintsch 412 used triply distilled water condensed and collected in quartz, but it was probably not completely gas-free when frozen; his data for k_e were given by graphs only.

82. MISCELLANEOUS ELECTRICAL DATA FOR ICE

Pyro-electric Effect.

J. Smithson ⁴²⁴ has observed that hail frequently consists of two hexagonal pyramids joined base to base. "One of the pyramids is truncated, which leads to the idea that ice becomes electrified on a variation of its temperature, like tourmaline, silicate of zinc, etc." This is the only mention of the probability of ice being pyroelectric that has come to the compiler's attention, though the observation of J. M. Adams, ⁴²⁵ that the ice-crystal is asymmetric with respect to its basal plane, indicates the same thing. Such asymmetry indicates the existence of piezo-electric properties also.

83. MAGNETIC SUSCEPTIBILITY OF ICE

Like water, ice is diamagnetic. G. Foex 426 has found that at the moment of freezing, the numerical value of the specific susceptibility (χ) decreases by 2.4 per cent. In their compilation, K. Honda, T. Ishiwara, T. Soné, and M. Yamada 427 give $\chi=-0.699$ micro-cgsm for the entire range 0 to $-120\,^{\circ}$ C, based on the observations of T. Ishiwara. 428 Taking the density of ice as 0.9168 g/cm³ at $0\,^{\circ}$ C, this gives for the volume susceptibility (κ) at $0\,^{\circ}$ C the value $\kappa=-0.641$ micro-cgsm.

More recently B. Cabrera and H. Fahlenbrach 429 have reported observations indicating that $\chi=-0.7019(1+0.000667t)$ micro-cgsm, and that the change on freezing is 2.2 per cent. This temperature coefficient, nearly 6 times that for water, is entirely incompatible with Ishiwara's observations, which extended to -120 °C, whereas Cabrera and Fahlenbrach did not go below -60 °C.

⁴²⁴ Smithson, J., Ann. Philos. (N. S.), 5, 340 (1823).

⁴²⁵ Adams, J. M., Proc. Roy. Soc. (London) (A), 128, 588-591 (1930) → Phys. Rev. (2), 36, 788 (A) (1930).

⁴²⁶ Foex, G., See Piccard, A., Arch. sci. phys. et nat. (4), 35, 209-231, 340-359, 458-482 (1913).

⁴²⁷ Honda, K., Ishiwara, T., Soné, T., and Yamada, M., Int. Crit. Tables, 6, 354-366 (356) (1929).

⁴²⁸ Ishiwara, T., Science Rep. Tohoku Univ., Sendai (1), 3, 303-319 (1914).

⁴⁰⁰ Cabrera, B., and Fahlenbrach, H., Am. Soc. Esp. Fis. y Quim., 31, 401-411 (1933).

III. Multiple-phase Systems

84. SURFACE-TENSION OF WATER

The number of articles treating of surface-tension and its measurement is very great, but in many cases, most unfortunately, the author of the article is not sufficiently acquainted with the mathematical derivation of the formula employed in obtaining the value of the surface-tension from the observed quantities to be able to appreciate its true significance. As a consequence, the experimental conditions realized by him frequently fail to accord with those demanded by the formula used, and the value of his discussion of the work, whether of himself or of another, is seriously impaired. Furthermore, and as a result of his failure to check its derivation, he occasionally uses a formula that is actually wrong, one involving a misprint or an algebraic error; and, in some cases, he merely guesses at the value of certain small corrections.

As a consequence, any mere assemblage of the various values published for the surface-tension of a given substance—such an assemblage as is commonly given in tables of constants—is of no assistance in enabling one to form an idea either of the most probable value of the surface-tension of that substance, or of the variability of its apparent surface-tension under good laboratory conditions, or of the possible dependence of its apparent surface-tension upon the method employed in measuring it.

Before such information can be obtained, each determination must be studied individually and in every detail, including the derivation of the formulas and their applicability to the experimental conditions actually realized. This involves great labor. In general, every determination based upon observations and computations that have been published without sufficient detail to enable one to make such a critical study should be summarily discarded as valueless; so should those for which the experimental conditions depart from those demanded by the formulas, unless the numerical value of the effect of such departure can be satisfactorily determined. I know of no publication of such a study of the existing data for surface-tension. Comparisons of selected groups of observations, of course, exist; and personal estimates of the most probable value of the surface tension of each of certain substances have been published from time to time. These estimates are presumably based upon some such detailed study as that just mentioned, but in some cases it is obvious that the study fell far short of what should be desired.

For values of the volumes of water menisci, see Tables 286 and 287.

Factors Possibly Affecting Surface-tension.

That the surface-tension of a liquid varies with the temperature and to a less extent with the nature of the overlying gas, and that its apparent value is greatly affected by even slight contamination of the surface, are well known facts. The first two are considered elsewhere, Tables 225 and 226, and text (p. 524). Of the third, nothing more need be said, as the data to be presented supposedly refer to uncontaminated gas-liquid surfaces.

But from time to time questions arise regarding the possible dependence of the observed surface-tension of a liquid upon other factors. They have to do with (1) the method used, (2) the material of the tube (in the method of capillary rise), (3) the effect of proximity to a solid wall, (4) the age of the (uncontaminated) surface, (5) the effect of illumination, (6) of electrification, (7) of a magnetic field, (8) of prolonged contact with catalysts. These, especially in their relation to water, will be considered here.

- (1). In comparing the results obtained by different methods, no effect arising either from the use of erroneous formulas or from a failure to secure the conditions demanded by the formulas used in the computations need be considered, for such effects result from mere blunders, and the data involved should be discarded unless the effects of the blunder can be eliminated. This greatly reduces the amount of data to be compared. Those left give no certain evidence of any difference that can unquestionably be attributed to a difference in the method, but a more careful comparison of the several available methods is much to be desired. It is entirely possible that the results obtained by dynamic methods may differ from those by static methods, and that the experimental details involved in some methods introduce unanticipated effects.
- (2). In 1894, P. Volkmann 1 found that the height to which water rises in a glass tube is independent of the nature of the tube; and recently, E. K. Carver and F. Hovorka 2 have found, contrary to the announced results of S. L. Bigelow and F. W. Hunter, 3 that the same is true for tubes of glass, zinc, copper, and silver.
- (3) It has long been recognized that the density, and even the structure, of a liquid in the immediate neighborhood of a solid may differ from that at a great distance from all solids. Any such difference would probably result in the tension of the gas-liquid surface near a solid wall being different from that elsewhere. In that case the form of the surface will differ from that corresponding to a surface of uniform tension, and, consequently, the value of the surface-tension as computed from the observations will differ from that for the surface far from solid walls, since all such computations are based on the assumption that the form of the surface is that corresponding to a surface of uniform tension. Such an effect

¹ Volkmann, P., Ann. d. Physik (Wied.), 53, 633-663 (1894).

Carver, E. K., and Hovorka, F., J. Am. Chem. Soc., 47, 1325-1328 (1925).
 Bigelow, S. L., and Hunter, F. W., J. Phys'l Chem., 15, 367-380 (1911).

might greatly exceed the dependence of the tension upon the nature of the solid. In the case of capillary rise, it would become of ever-increasing importance as the diameter of the tube is reduced.

W. A. Patrick and N. F. Ebermann 4 have published observations which they interpret as indicating that the pressure of the vapor in equilibrium with the concave liquid-gas surface in a tube of very small diameter (a few microns) is less than that computed for a surface of the same curvature and the tension characteristic of a flat surface of the liquid far from solid walls (for formula, see p. 568). This suggests that the tension is greater for the smaller surface, presumably on account of the presence of the solid walls. But the actual significance of their observations is not entirely clear, and the interpretation of them is correspondingly difficult. In at least some cases, the interpretations offered may overlook essential factors, as pointed out by D. J. Woodland and E. Mack, Jr. That the observations are not to be accounted for by the mere curvature of the surface is indicated by the earlier observation of N. Gudris and L. Kalikowa 6 that the partial pressure of the vapor in equilibrium with air-suspended water droplets 0.1 to 1.0 μ in diameter is equal to that computed by Kelvin's (Thomson's) formula (p. 568). Furthermore K. W. v. Nägeli had found that the pressure required to drive water from tubes 3 to 9μ in diameter is about the same as would be inferred from observations on much larger tubes. He used freshly drawn tubes, and stated that older tubes always gave off air, forming minute bubbles in the water, which impeded the flow.

The subject is intimately connected with the least thickness of an adsorbed layer that has the same vapor pressure as does the liquid in bulk, and with the least thickness of a layer of water that exhibits true viscosity and has a viscosity that is the same as that of the liquid in bulk, although it is not identical with either of these. L. J. Briggs 8 studied the adsorbed layer of water on quartz when in equilibrium with atmospheres of various relative humidities at 30 °C. He found that at 99 per cent humidity the amount of that adsorbed layer that could be removed by heating to 110 °C corresponded to a thickness of 0.027 μ. Similarly, I. R. McHaffie and S. Lenher 9 observed that the vapor-pressure of adsorbed water films varied with the thickness unless that exceeded several hundreds of molecules if on glass, and several tens of molecules if on platinum (300 molecules = 0.09 μ , approximately). And S. H. Bastow and F. P. Bowden 10, 11 have observed that even at 0.1 °C a film of water only 0.2 u thick, flowing

⁴ Patrick, W. A., and Ebermann, N. F., Idem. 29, 220-228 (1925). Cf. Shereshefsky, J. L., J. Am. Chem. Soc., 50, 2966-2980, 2980-2985 (1928); Latham, G. H., Idem, 50, 2987-2997 (1928). 8 Woodland, D. J., and Mack, E., Jr., J. Am. Chem. Soc., 55, 3149-3161 (1933). ⁶ Gudris, N., and Kalikowa, L., Z. Physik, 25, 121-132 (1924).

⁷v. Nägeli, K W, Sitz.-ber. Bayer Akad. Wiss. München, 1866, 353-376 (358) (1866).

⁸ Briggs, L. J., J. Phys'l Chem., 9, 617-640 (1905).

⁹ McHaffie, I. R., and Lenher, S., J. Chem. Soc. (London), 127, 1559-1572 (1925).

¹⁰ Bastow, S. H., and Bowden, F. P., Proc. Roy. Soc. (London) (A), 151, 220-233 (1935);

Cf. Idem, 134, 404-413 (1932). 11 Bowden, F. P., and Bastow, S. H., Nature, 135, 828 (L) (1935).

Table 225.—Surface-tension of Water

(For thermal variations, see also Table 226.)

All values at temperatures not exceeding 100 °C refer to an air-water surface at atmospheric pressure, the water being, presumably, saturated with air, and the air with water-vapor. Those above 100 °C refer to the surface separating pure water from its pure vapor. For the effect of a change in the gas, see Table 229.

As primary standard for the ICT values, Young and Harkins accepted for the tension of a water-air surface at 20 °C the value 72.75 ± 0.05 dyne/cm, which they derived from the determinations of T. W. Richards and L. B. Coombs ²⁹ which were corrected by T. W. Richards and E. K. Carver ³⁰ to yield 72.72; of W. D. Harkins and F. E. Brown ³¹ giving 72.80; of T. W. Richards and E. K. Carver ³⁰ giving 72.73; and of T. F. Young and P. L. K. Gross (unpublished) giving 72.80, all determined by the rise in capillary tubes.³²

At the critical point the surface-tension does not become zero when the meniscus vanishes.³³

In contrast to the other values tabulated below, the TB ones show an anomaly near 13 °C.

For sea-water containing s grams of salts per kg, $\gamma_s = \gamma_o + 0.0221s$ dyne/cm, where γ_s and γ_o are the air-liquid surface-tensions of the seawater and of pure water, respectively, both at the same temperature.³⁴ If s = 35, the average amount of salts, then $\gamma_s - \gamma_o = 0.77$ dyne/cm.

 $\gamma=$ surface-tension; $a^2=2\gamma/(\rho-\sigma)g$; g= acceleration of gravity (here taken as 980.665 cm/sec²); $\rho=$ density of liquid; $\sigma=$ density of the adjacent gas (here taken, the gas being air, as 0.001200(293.1)/(273.1+t)=0.3517/(273.1+t) g/cm³, the temperature being t °C); $\Delta\equiv\gamma-\gamma_c$ where $\gamma_c=75.64-0.13910t-0.0003000t²$ dyne/cm is an empirical formula which represents the ICT values fairly well; τ is the tolerance assigned by Young and Harkins.

Unit of γ , τ , and $\Delta=1$ dyne/cm = 1 gram sec⁻² = 0.10197 mg*/mm; of $a^2=1$ cm² = 100 mm². Temp. = t °C

Refa _		1	СТ		M	oser	War	ren	тв
t	γ	τ	a^2	70	γ	100Δ	γ	100Δ	100Δ
$ \begin{array}{r} -8 \\ -5 \\ 0 \\ +5 \end{array} $	76.9 ₆	0.3	0.1574	76.734					
5	76.42	0.2	0.1562	76.32_{8}					
0	75.64	0.1	0.1544_{s}	75.64 ₀	75.62°	-2	75.94	+30	+36
+5	74.9 ₂	0.1	0.1529_{9}	74.93 ₆	74.86	-8	75.19	+25	-44
10	74.22	0.05	0.1516_{0}	74.21 _e	74.12	-10	74.43	+21	-72
11	74.07	0.05	0.1513_{1}	74.07	73.96	-11			
12	73.93	0.05	0.1510_{3}	73.92 ₈	73.80	-13			
13	73.78	0.05	0.1507₅	73.781	73.65	-13			
14	73.64	0.05	0.1504_{8}	73.634	73.50	-13			
15	73.49°	0.05	0.1501_{9}	73.48 ₆	73.35	-14	73.65	+16	-37
16	73.34	0.05	0.1499_{1}	73.337	73.20	-14			
17	<i>7</i> 3.19	0.05	0.1496_{3}	73.18 ₈	73.04	-15			
18	73.05	0.05	0.1493_{7}	73.03 ₉	72.89	-15			
19	72.90	0.05	$0.1490_{\rm p}$	72.88 ₀	72.73	-16			
20	72.75	0.05	0.14881	72.73 ₈	72.58	-16	72.86	+12	+13

Table 225.—(Continued)

Ref.a		I	CT		Moser		Warren		TB
t	γ	•	α^2	70	γ	100∆	γ	100∆	100∆
21	72.59	0.05	0.1485_{2}	72.587	72.43	-16			
22	72.44	0.05	0.14824	72.435	72.27	-16			
23	72.28	0.05	0.1479 ₆	72.282	72.10	-18			
24	72.13	0.05	0.1476_{8}	72.12	71.96	-17			
25	71.97	0.05	0.1473_{8}	71.974	71.81	-16	72.09	+12	+3
26	71.82	0.05	0.1471_{1}	71.820	71.65	-17			
27	71.66	0.05	0.1468_{a}	71.66	71.50	-16			
28	71.50	0.05	0.14654	71.51.	71.34	-17			
29	71.35	0.05	0.1462_{7}	71.354	71.19	-16			
30	71.18	0.05	0.1459_{7}	71.19_{7}	71.03	-17	71.33	+13	+5
35	70.38	0.05	0.1445_{6}	70.40_{7}	70.23	-18	70.54	+13	-4
40	69.56	0.05	0.1431_{8}	69.59 _e	69.42	-18	69.73	+13	-10
45	68.74	0.05	0.1417_{8}	68.772	68.59	-18	68.83	+6	
50	67.91	0.05	0.1403_{2}	67.93 ₅	67.75	-18	68.02	+9	
55	67.05	0.05	0.1388_{7}	67.082	66.84	-24	67.14	+6	
60	66.19	0.05	0.1374_{1}	66.21,	66.04	-17	66.24	+3	
70	64.42	0.1	0.1344_{9}	64.43 ₃	64.28	-15	64.51	+8	
80	62.61	0.1	0.1315	62.592	62.50	-9	62.69	+10	
90	60.7 _b	0.2	0.1284	60.69_{1}	60.68	-1	60.80	+11	
100	58.8 ₅	0.2	0.1253	58.73 _o	58.80	+7			
110	56.8°	0.2		56.70 ₀					
120	54.8 ₀ d	0.2		54.628					
130	52.84d	0.3		52.487					

References:

From compilation of T. F. Young and W. Harkins, 35 based upon the accepted value at 20 °C (see head of table) and the observations of C. Brunner, 36 W. I. Humphreys and J. F. Mohler, 37 J. L. R. Morgan and C. E. Davis, 36 J. L. R. Morgan and A. McD. McAfee, 39 W. Ramsay and J. Shields, 40 T. W. Richards, C. L. Speyers, and E. K. Carver, 41 H. Sentis, 42 S. Sugden, 43 and P. Volkmann. 44 Moser, H., Ann. d. Physik (4), 82, 993-1013 (1927).

Timmermans, J., and Bodson, H., Compt. rend., 204, 1804-1807 (1937); values were read from their graph.

Warren Warren, E. L., Phil. Mag. (7), 4, 358-386 (1927).

 6 For water at 0 °C, G. Schwenker 45 has found $\gamma = 75.59_{7}$ with an estimated uncertainty not exceeding 0.044 per cent.

° By a method involving an impact of two jets, W. N. Bond 60 found for a water-air surface that was renewed about 80 times a second, $\gamma = 73.83 \pm 0.13$ dyne/cm at 15 °C. 6 These values are for water in contact with its own pure vapor.

¹⁸ Bulkley, R., Bur. Stand J. Res., 6, 89-112 (RP264) (1931).

¹⁸ Bohr, N., Phil. Trans. (A), 209, 281-317 (1909).

¹⁴ Hiss, R., Diss., Heidelberg, 1913.

¹⁵ Schmidt, F., and Steyer, H., Ann. d. Physik. (4), 79, 442-464 (1926).

¹⁶ Kleinmann, E., Idem, 80, 245-260 (1926).

¹⁷ Young, T. F., and Harkins, W. D., Int. Crit. Tables, 4, 446-475 (474) (1928).

¹⁸ Lenard, P., Sitz. Heidelberger Akad. Wiss. (A), 5, No. 28, pp. 16-23 (1914).

¹⁹ Seitz, E. O., Ann. d. Physik (5), 1, 1099-1108 (1929).

²⁰ Buchwald, E., and König, H., Idem, 23, 557-569 (1935).

²¹ Grumbach, A., and Schlivitch, S., Compt. rend., 181, 241-243 (1925).

²² Auer, H., Z. Physik, 66, 224-228 (1930).

²⁰ Johner, W., Helv. Phys. Acta, 4, 238-280 (1931) = Diss., Bern, 1930.

²⁴ Piccard, A., Arch. Sci. phys. et nat. (4), 35, 209-231, 340-359, 458-482 (1913).

²⁵ Liebknecht, O., and Wills, A. P., Ann. d. Physik (4), 1, 178-188 (1900).

²⁶ Quincke, G., Idem (Pogg.), 160, 560-588 (586) (1877); Idem (Wied.), 24, 347-416 (376-377) (1885).

²⁷ Brunner, C., Idem (Pogg.), 79, 141-144 (1850) reporting observations of J. R. A. Mousson.

²⁸ Baker, H. B., J. Chem. Soc. (London), 1927, 949-958 (1927).

Richards, T. W., and Coombs, L. B., J. Am. Chem. Soc., 37, 1656-1676 (1915).

⁸⁰ Richards, T. W., and Carver, E. K., Idem, 43, 827-847 (1921).

Table 226.—Thermal Variation of the Surface-tension of Water

For the value of the surface-tension at each of a selected number of temperatures, see Table 225. For the values of the molecular surface energy at various temperatures, see Table 227.

In this table are given for the air-water surface the values of: I. The temperature derivative $(d\gamma_e/dt)$ as defined by those empirical equations of the form $\gamma_o = \gamma_0(1-at-bt^2) = \gamma_0(1-\delta)$ that have been found to represent, respectively, the corresponding sets of data given in Table 225 (the TB data are not here included, having appeared after the completion of this table). II. The several values of $\delta (= (\gamma_0 - \gamma_e)/\gamma_0)$ corresponding to the same three equations and to others of the same form that may be found in other compilations. III. The several values of δ corresponding to certain other types of interpolation equations.

Unit of $\gamma = 1$ dyne/cm; of $d\gamma/dt = 1$ dyne-cm⁻¹ per 1 °C. Temp. = t °C

I. Temperature derivative $d\gamma_e/dt$. $\gamma_e = \gamma_0(1 - at - bt^2)$. Dr. Domke ⁴⁷ concluded that the data available in 1902 indicated that at 20 °C $d\gamma/dt = -0.151$.

Ref.a→	ICT	M	Wa	Ref.a→	ICT	M	Wa
1000a→	1.83 _p	1.95₀	1.91,	$1000a \rightarrow$	1.83_{0}	1.95	1.91,
10°b→	3.97	2.62	3.33	10°b→	3.97	2.62	3.33
$\gamma_0 \rightarrow$	75.64	75.62	75.91	$\gamma_0 \rightarrow$	75.64	75.62	75.91
t		$-d\gamma_o/dt$		t		$-d\gamma_e/dt$	
5	0.136	(0.146)	(0.143)	45	0.166	0.166	0.168
0	0.139	0.148	0.145	50	0.169	0.168	0.170
+5	0.142	0.150	0.148	55	0.172	0.170	0.173
10	0.145	0.152	0.150	60	0.175	0.172	0.176
15	0.148	0.154	0.153	70	0.181	0.176	0.180
20	0.151	0.156	0.155	80	0.187	0.180	0.186
25	0.154	0.158	0.158	90	0.193	0.184	0.191
30	0.157	0.160	0.160	100	0.199	0.188	0.196
35	0.160	0.162	0.163	110	0.205	(0.192)	(0.201)
40	0.163	0.164	0.165	120	0.211	(0.196)	(0.206)

II. Values of $\delta \equiv (\gamma_0 - \gamma_e)/\gamma_0$; $\gamma_e = \gamma_0(1 - at - bt^2)$.

Ref.a→	ICT	M	Wa	\mathbf{F}	S	RSC
1000a→	1.83 ₀	1.95_{0}	1.91,	1.90_{2}	2.02	2.08_{0}
10°b.→	3.97	2.62	3.33	2.50	0	3.29
$\gamma_{o} \rightarrow$	75.64	75.62	75 .91			75.89
t			100δ			
-10	-1.80	(-1.9_3)	(-1.9)	(-1.9)	-2.0	-2.1
-5	-0.9_{1}	(-0.9_7)	(-1.0)	(-0.9)	-1.0	-1.0
0	0	0	0	` 0 ´	0	0
$^{0}_{+5}$	+0.9s	$+0.9_{0}$	+1.0	+1.0	+1.0	+1.0
10	$1.8_{\rm s}$	1.9_{0}	1.9	1.9	+2.0	2.1
20	3.84	4.02	4.0	3.9	4.1	4.3
25	4.8_{5}	5.0_{6}	5.0	4.9	5.1	5.4
30	5.8 ₈	6.12	6.0	5.9	6.1	6.6
40	8.0_{0}	8.26	8.2	(8.0)	8.1	8.9
60	12.47	12.6 _e	12.7	(12.3)	12.2	13.7
80	17.2_{5}	17.3	17.4	(16.8)	16.2	18.8
100	22.3	22.31	(22.4)	(21.5)	20.3	24.2
120	27.81	(27.2_{0})	(27.7)	(26.4)	24.3	29.8

Table 226,-(Continued)

III. Richards, Speyer, and Carver 41 have proposed also the formula $\gamma_e = K(1 - T/T_o)^{\alpha}$, with $\alpha = 1.2$; T and T_o are, respectively, the absolute temperatures corresponding to ye and to the critical point of water, K is a constant. That value of α is obviously incorrect (cf. columns 3 and 4); if they recorded the cotangent instead of the tangent of the slope of the logarithmic graph, then α should be 1/1.2 = 0.83; that value leads to the values in column 5. They represent the RSC data fairly well. The ICT data cannot be satisfactorily represented by an equation of that form, the logarithmic graph being curved; they are, however, fairly well represented by the formula $\gamma_e = K(1 + ct + dt^2)(1 - T/T_c)^{0.849}$. For convenience we may write $(1 + ct + dt^2) \equiv 1 + \epsilon = f(t)$. $(\equiv \delta) = 1 - f(t)(1 - t/t_c)^{0.849}$ where t and t_c are the centigrade temperatures corresponding to T and T_c , respectively; $t_c = 374.0 \,^{\circ}\text{C}$; $T_c =$ 647.1 °K.

Weinstein (We) has proposed the formula $\gamma_{\theta} = 73.49(1 - 0.001458t)\rho$ dynes/cm; whence we find $\delta (\equiv (\gamma_0 - \gamma_e)/\gamma_0) = 1 - (1 - 0.001458t)\rho/\rho_0$.

In the following tabulation, experimental data taken from the preceding section of this table are given in columns 2 and 3, and in the other columns are values computed by the formulas just given. It is obvious that only column 6 accords satisfactorily with 2; the values in 4 and 8 are entirely unsatisfactory.

1 Ref.a→	2 ICT	3 RSC	4 RSC	5	6	7	8 We
$\alpha \rightarrow t$			1.2 100δ	0.83	0.849	100€ ^b	100δ
$-10 \\ -5$	-1.80 -0.91	-21 -1.0	-3.2 -1.6	-2.2 -1.1	−1.7 −0.9	-0.55 -0.27	-0.7
$^{0}_{+5}$	$^{0}_{+0.93}$	$^{0}_{+1.0}$	$^{0}_{+16}$	$^{0}_{+1.1}$	$^{0}_{+0.9}$	$^{0}_{+0.25}$	$^{0}_{+0.7}$
10 20	1.88 3.84	2.1 4.3	3.2 6.4	2.2 4.5	1.8 3.7	0.48 0.88	1.5 3.1
25 30	4.85 5.88	5.4 6.6	8.0 9.5	5.6 6.7	4.7 5.7	1.04 1.20	3.9 4.8
40 60	8.00 12.47	8.9 13.7	12.7 18.9	9.0 13.6	7.8 12.3	1.44 1.70	6.6 10.3
80	17.25	18.8	25.1	18.2	17.1 22.2	1.64 1.28	14.1 18.1
100 120	22.36 27.81	24.2 29.8	31.1 37.1	22.8 27.6	27.6	0.60	22.1

a References:

Forch, C., Ann. d. Physik (4), 17, 744-762 (1905). See Table 225.

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M

Moser, H., Ann. d. Physik (4), 82, 993-1013 (1927). Richards, T. W., Speyer, C. L., and Carver, E. K.⁴¹ Sentis, H., Jour. de Phys. (3), 6, 183-187 (1897); Ann. Univ. Grenoble, 27, 593-624 (1915). RSC

Warren, E. L., Phil. Mag. (7), 4, 358-386 (1927). Wa

Weinstein, B., Metron. Beitr. (Norm. Aich. Komm.), No. 6 (1889).

^b From the formula $\gamma_e = K(1+\epsilon)(1-T/T_e)^{0.849}$; $\epsilon \equiv 0.000515t - 0.00000387_bt^2$.

between solid walls, has a true viscosity that is, within experimental error (say 10 per cent), the same as for water in bulk.

Furthermore, in the experiments of R. Bulkley, 12 on the viscous flow of liquids through very fine capillaries, the effect of the walls in modifying the pertinent properties of the liquid was inappreciable, although he could have detected it had it been equivalent to the production of a stationary film only 0.03 µ thick.

All these indicate that the effect of the walls extends no farther than a very small fraction of a micron, and, consequently, will not affect the surface-tension at more distant points.

See also p. 527.

(4). In 1909, N. Bohr ¹³ concluded that the tension of a clean gas-liquid surface does not change after it is 0.06 sec old. More recently, R. Hiss, 14 F. Schmidt and H. Steyer, 15 and E. Kleinmann 16 have investigated younger surfaces, seeking for evidence that the tension of a newly formed surface exceeds that of an equally clean surface of greater age. All used the same general method, and found an apparent progressive decrease in the tension, equilibrium being reached after a few milliseconds. Whether the effect observed was due to an actual decrease in the tension or was a secondary phenomenon due to an unsatisfactory technique is not entirely clear. T. F. Young and W. D. Harkins 17 seem inclined to ascribe it to the latter. P. Lenard ¹⁸ has given reasons for expecting the tension to decrease as the surface ages.

More recently, the subject has been studied by E. O. Seitz ¹⁹ and by E. Buchwald and H. König.²⁰ The latter used a novel method. Each found that the surface-tension decreased as the surface aged, the decrease being approximately exponential.

(5). A. Grumbach and S. Schlivitch ²¹ report that illuminating a vapor-water surface does not change its tension. The tension of certain other liquid surfaces is changed by illumination.

(6). The few reported observations on the tension of electrified gaswater surfaces give no indication of any effect of electrification, other than those resulting from electrostatic repulsion.

(7). No one has succeeded in showing that a magnetic field has any effect upon the surface tension. H. Auer 22 has recently reported that the application of a horizontal magnetic field of 20,000 gauss to the air-water meniscus in a vertical capillary does not change the surface tension by as

⁸¹ Harkins, W. D., and Brown, F. E., Idem, 41, 499-524 (1919). ⁸² Cf. Lenard, P., v. Dallwitz-Wegener, R., and Zachmann, E., Ann. d. Physik (4), 74, 381-404 (1924).

⁸⁸ Winkler, C. A., and Maass, O., Can. J. Res., 9, 65-79 (1933).

⁸⁴ Krümmel, O., "Handbuch der Ozceanographie," Vol. 1, pp. 280-281, 1907. Krummel, C., "Handbuch der Ozceanographie," Vol. 1, pp. 280-281, 1907.
 Young, T. F., and Harkins, W., Int. Crit. Tables, 4, 446-475 (447) (1928).
 Brunner, C., Ann. d. Physik (Poag.), 70, 481-529 (1847).
 Humphreys, W. J., and Mohler, J. F., Phys. Rev., 2, 387-391 (1895).
 Morgan, J. L. R., and Davis, C. E., J. Am. Chem. Soc., 38, 555-568 (1916).
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 Ramsay, W., and Shields, J., J. Chem. Soc. (London), 63, 1089-1109 (1893).
 Billeds, T. W. Species, C. L. and Carver, F. K. J. Am. Chem. Soc., 46, 111

⁴ Richards, T. W., Speyers, C. L., and Carver, E. K., J. Am. Chem. Soc., 46, 1196-1207 (1924).

much as 10 parts in a million. He varied the temperature from 15 to 92 °C. See also, W. Johner, 28 A. Piccard, 24 O. Liebknecht and A. P. Wills,25 G. Quincke,26 and C. Brunner.27

(8). It has been stated that prolonged contact of water with such catalysts as charcoal, thoria, and platinum so modifies it that its surfacetension is increased.28

Molecular Surface Energy.

The molecular surface energy is $\gamma_M = \gamma (M/\rho)^{2/3}$ by definition, γ being the surface-tension of the liquid in contact with its own pure vapor, M the molecular weight of the vapor, and ρ the density of the liquid. It is the mechanical work required to increase the area of the surface by an amount equal to the area of one face of the cube of liquid of mass equal to one mole of the vapor, the temperature being maintained constant by a suitable addition of heat.

The number of molecules contained in such a cube of liquid is the same for every liquid having the same fixed ratio between its molecular weight and the molecular weight of its vapor; and the number of molecules contained in the surface layer of area equal to the face of that cube and of thickness equal to a fixed multiple of the mean distance between adjacent molecules is likewise the same for every such liquid, provided the variation in the mean distance between adjacent molecules as the surface is approached is the same function of the distance from the surface, all distances being expressed in terms of the mean distance between adjacent molecules in the body of the liquid; i.e., in terms of the length of an edge of the cube. In particular, if the molecular weight of each liquid is the same as that of its vapor, and if the mean distance between adjacent molecules does not change as the surface is approached, then each such cube contains one mole of liquid, and each surface layer of area equal to one face of the cube and of thickness equal to a fixed multiple of the mean distance between adjacent molecules (i.e., of thickness proportional to an edge of the cube) will contain the same number of molecules of the liquid; and enlarging the liquid surface by an area equal to the face of the cube will bring the same number of molecules from the interior of the liquid into the surface layer, whatever the liquid may be.

It is for these reasons that M/ρ may be called the molecular volume, $(M/\rho)^{2/8}$ the molecular area, and $(M/\rho)^{1/3}$ the molecular distance. Their molecular significance is independent of the nature of the substance. For the same reasons it seemed probable that fundamental relations would be discovered more readily by studying γ_M rather than γ .

⁴⁹ Sentis, H., Ann. Univ. Grenoble, 27, 593-624 (1915).

⁴⁸ Sugden, S., J. Chem. Soc. (London), 119, 1483-1492 (1921); 121, 858-866 (1922).

⁴⁴ Volkmann, P., Ann. d. Physik (Wied.), 56, 457-491 (1895). 45 Schwenker, G., Ann. d. Physik (5), 11, 525-557 (1931). 40 Bond, W. N., Proc. Phys. Soc. (London), 47, 549-558 (1935).

⁴⁷ Domke, Wiss. Abh. Norm.-Aich. Komm., 3, 1-99 (1902).

It will be noticed that, contrary to the implications of statements to be found in certain texts, neither the conception of molecular surface energy (γ_M) nor the definition of $(M/\rho)^{2/3}$ as the molecular area has anything directly to do with a sphere.

In 1886, R. Eötvös ⁴⁸ propounded a number of relations based upon supposed analogies and the idea of corresponding states, culminating in the conclusion that $d\gamma_M/dt$ is a constant, independent of the nature of the substance and of the temperature. This relation he tested by means of existing data, and found it to be satisfied in many cases, the value of the derivative being quite close to -2.12 ergs per gram-molecular area and per 1 °C. Whence the negative derivative has been called the Eötvös constant; and is often denoted by k_B ; $k_B \equiv -d\gamma_M/dt$. He appears to have made no attempt to test the validity of the intermediate relations upon which this final one was based, and W. Ramsay and J. Shields ⁴⁹ showed that they did not accord with observations.

Inspired by the work of Eotvös, but starting from a different set of supposed analogies, Ramsay and Shields ⁴⁹ reached the conclusion that γ_M should be proportional to $(t_o - t)$; that is, to the temperature measured downward from the critical temperature. But in order to secure agreement with experimental data, it was necessary to decrease that temperature by an empirically determined amount (θ) , usually about 5 or 6 °C, and then, in order to make $\gamma_M = 0$ when $t = t_o$ they introduced an exponential term, giving finally the relation $\gamma_M = k_B \left[t_o - t - \theta (1 - 10^{-\lambda(t_o - t)}) \right]$. Except for the exponential term, which is negligible unless $(t_o - t) < 30$ °C, this leads at once to the relation found by Eötvös; it is merely a special case of the integral of his relation.

They confirmed the conclusion of Eötvös, that k_B has essentially the same constant value for each of many substances, the average being 2.12₁. For such substances, they assumed that the molecular weight is the same in the liquid as in the vapor phase.

But for water and certain other substances k_B is much less than 2.12, and varies with the temperature, increasing as the temperature rises. This they ascribed to an association of some of the molecules, causing the effective molecular weight of the liquid to be xM, where x, called by them the constant of association, is greater than unity. Then, for $(t_0 - t) > 30$ °C, their relation becomes $x^{2/3}\gamma_M = k [t_0 - t - \theta]$, where $k = 2.12_1$.

Whence,
$$x^{2/3} d\gamma_{M}/dt + (2/3)x^{-1/3}\gamma_{M} \frac{dx}{dt} = -k$$
, k_{E} ($\equiv -d\gamma_{M}/dt$) =

 $kx^{-2/8} + (2/3)(\gamma_M/x)dx/dt$, and $x^{2/8} = k \div [k_B - (2/3)(\gamma_M/x)dx/dt]$. As dx/dt is negative, the two terms in the right-hand member of the second expression conspire to make $k_B < k$. The determination of x, dx/dt, and θ in terms of k, γ_M , and k_B is, in general, difficult, if not impossible. But

⁴⁸ Eotvos, R., Ann. d. Physik (Wied.), 27, 448-459 (1886).

⁴⁰ Ramsay, W., and Shields, J., Phil. Trans. (A), 184, 647-674 (1893).

⁵⁰ Ramsay, W., and Shields, J., J. Chem. Soc. (London), 63, 1089-1109 (1893).

Ramsay ⁵¹ observed that, at least in certain cases, γ_{M} can be expressed in the form $\gamma_{M} = k'(t_{o} - t - \theta)/[1 + \mu(t_{o} - t)]$. If it is assumed that this θ is the same as the θ in the relation $\gamma_{M}x^{2/3} = 2.12(t_{o} - t - \theta)$, then $x^{2/3} = 2.12[1 + \mu(t_{o} - t)]/k'$. This is Ramsay's procedure, and has been used in deriving the values of x_{R} given in Table 227. For the ICT data

Table 227.-Molecular Surface Energy of Water

See text for references, discussion of the significance of the several quantities, etc.

 $\gamma_{M} \equiv \gamma(M/\rho)^{2/3}$; $k_{E} \equiv -d\gamma_{M}/dt$; k_{E0} = value of k_{E} tabulated in ICT; M=18.0154, the formula-weight of $H_{2}O$; $\rho=$ density of water as given in Tables 93 and 255; $\gamma=$ tension of the water-air, or of the water-vapor, surface, the latter for the Ramsay data and for the ICT data above $100\,^{\circ}\text{C}$; $\Delta_{10} \equiv (\gamma_{M})_{t} - (\gamma_{M})_{t+10}$; $x_{a} \equiv (2.12_{1}/k_{E})^{1.5}$; $x_{am} = (21.2_{1}/\Delta_{10})^{1.5}$; $x_{E} = \text{association constant computed by Ramsay's method (it is of little value, see text). In computing the ICT data, the values called <math>\gamma_{0}$ in Table 225 have been used; in computing the Ramsay data, his observed values of γ were used (they are all much too small); k_{E} has been computed from the values of γ and ρ , and of their variations with t, by means of the relation given in the text.

Unit of $\gamma_{\underline{M}}=1$ erg per g-molecular area; of $k_{\underline{B}}=1$ erg per g-molecular area, per 1 °C; of $(M/p)^{2/3}=1$ cm²; κ_a , κ_a , and $\kappa_{\underline{B}}$ are pure numbers. Temp. = t °C.

				· ICT —					- Ramsa	y -		
*	$k R_0$	k H	Xa	γ_M	Δ_{10}	Xam	x_R	γ_{M}	Δ_{10}	xam	x_R	$(M/\rho)^{2/8}$
0		0.980	3.17	519.86	9.72	3.22	1.46	503.16	8.69	3.8	1.53	6.8728
10		0.968	3.24	510.14	9.68	3.25	1.43	494.47	8.73	3.8	1.50	6.8734
20		0.972	3.22	500.46	9.77	3.19	1.40	485.74	9.50	3.3	1.47	6.8803
25	$1.0_{\rm a}$	0.977	3.20	495.58								6.8856
30		0.984	3.15	490.69	9.93	3.12	1.37	476.24	9.96	3.1	1.44	6.8921
40		1.004	3.06	480.76	10.15	3.03	1.34	466.28	9.21	3.3	1.41	6.9079
50		1.029	2.96	470.61	10.42	2.90	1.31	457.07	10.39	2.9	1.38	6.9274
60		1.055	2.85	460.19	10.72	2.78	1.28	446.68	10.34	2.9	1.35	6.9501
70	1.0_{7}	1.085	2.73	449.47	11.05	2.66	1.25	436.34	10.19	3.0	1.32	6.9758
80		1.125	2.59	438.42	11.41	2.53	1.22	426.15	11.61	2.5	1.29	7.0044
90		1.160	2.46	427.01	11.80	2.41	1.19	414.54	11.50	2.5	1.27	7.0357
100	1.1_{8}	1.200	2.35	415.21	12.36	2.24	1.16	404.04	11.56	2.5	1.24	7.0698
110		1.248	2,22	402.85	12.76	2.14	1.13	392,48	11.87	2.4	1.21	7.1038
120	1.2_{7}	1.297	2.08	390.09	13.21	2.03	1.10	380.61	11.25	2.6	1.18	7.1409
130	•	1.344	1.98	376.88			1.09	369.36			1.15	7.1805

for water, k'=3.618 ergs per g-molecular area, per 1 °C, $\theta=58.2$ °C, and $\mu=0.00321$ (°C)⁻¹; for the Ramsay data for water, k'=2.994 ergs per g-molecular area, per 1 °C, $\theta=43.7$ °C, and $\mu=0.00258$ per 1 °C. In each case $t_o=374.0$ °C. The constants for the Ramsay data have been completely recalculated, as his calculation was based upon $t_o=358.1$ °C, and upon densities that differ slightly from those in Table 93 and those derivable from Table 255.

⁵¹ Ramsay, W., Proc. Roy. Soc. (London), 56, 171-182 (1894) = Z. physik Chem., 15, 106-116 (1894).

It should, however, be realized that Ramsay's procedure is very arbitrary. There is no reason whatever for believing that the two θ 's must have the same value. Rather the contrary; for the empirically determined value of the θ in the expression for γ_M for water is about 10 times as great as the value found for substances for which $k_{\rm E}=2.12$. (Ramsay found a value only about 4 times as great, but that was because he used too low a value for the critical temperature.) Furthermore, the value (x_R) so found for the association constant of water is much less than 2, while we now have other reasons for believing that the association constant of water is at least as great as 2. Consequently, little, if any, weight should be attached to the actual, numerical values of x_R . I doubt if they are, in general, worth the time required to compute them. They seem to be of no more value than the much more readily computed quantities $x_a \equiv (2.12_1/k_E)^{1.5}$ and $x_{am} \equiv \{2.12_{17}/(\gamma_{M,t} - \gamma_{M,t+\tau})\}^{1.5}$, which may be called the coefficients (actual and mean, respectively) of apparent association. If the Eötvös-Ramsay relation applied and x were independent of t, then $x_a = x$; while if x varied with t, x_{am} would be the mean value of x over the range t to $t + \tau$. Also at those high temperatures, if such exist, at which x_a and x_{am} are independent of t, $x_a = x_{am}$ and each is actually the coefficient of association as defined by the Eötvös-Ramsay relation. The value of $k_R \equiv$ $-d\gamma_M/dt$ may be determined either directly from a formula expressing the variation of γ_M with t, or by means of the relation $-d\gamma_M/dt = (M/\rho)^{2/3}$ $\{(2/3)(\gamma/\rho)(d\rho/dt)-d\gamma/dt\}.$

A theoretical discussion of the surface energy of liquids has recently been published by H. Margenau.⁵²

Angle of Contact.

The angle of contact (the angle including the liquid) between an airwater, or a vapor-water, surface and a glass surface covered with a film of water is zero.⁵³

In general, the contact angle between a gas-liquid surface and any solid covered by a film of the liquid is zero. When the solid is not covered by a film of the liquid, the angle of contact is, in general, variable, the line of contact exhibiting a reluctance to move over the solid. (For a suggested explanation, see *Miscellanea*—p. 526.) In such cases, the characteristic angle of contact is commonly taken as the mean of the greatest and the least equilibrial angle that can be obtained; the former occurs when equilibrium has been attained at the end of an advance of the line of contact toward the uncovered ("dry") portion of the solid; the latter, at the end of an advance in the opposite direction. Angles computed from measurements made on stationary sessile drops are of little value unless they are the means of such greatest and least values.

Margenau, H., Phys. Rev. (2), 38, 365-371 (1931).
 Young, T. F., and Harkins, W. D., Int. Crit. Tables, 4, 434 (1928), on the strength of the observations of: Anderson, A., and Bowen, J. E., Phil. Mag. (6), 31, 143-148 (1916); Bosanquet, C. H., and Hartley, H., Idem, 42, 456-461 (1921); Richards, T. W., and Carver, E. K., J. Am. Chem. Soc., 43, 827-847 (1921); Sentis, H., Jour. de Phys. (3), 6, 183-187 (1897).

It seems that the contact angle is sensitive to changes in the structure of the solid, F. E. Bartell, J. L. Culbertson, and M. A. Miller 54 having found that it has a relatively large value (30 to 80°) for Pyrex and silica when there are great internal strains, but a zero value when the solids have been well annealed. They found also that the value of the angle for brass, as well as for Pyrex, can be changed by compressing the solid. See also Bartell and Miller.55

Table 228.—Contact Angle between Air-water Surfaces and Dry Solids

 θ is the angle containing the limiting wedge of liquid.

If a solid is wet, i.e., is thoroughly covered with a layer of water, then $\theta=0$.

	Unit of $\theta = 1^{\circ}$ of arc.	Temp. = t °C		
	Solid	t	θ	Ref.
			68 ± 4	BM
Platinum			63 ± 4	BM
Copper sulfide			0 ,	DW
Azobenzene		14	77	BH, ICT, M
Apple wax			74	M
Paraffin		14	106.7	BH, ICT
			104.6	A
Paraffin			105	TL, KB
Paraffin			110 ± 6	
Talc		25	86	BZ
Glyptol resin ("Glypto	01 1350")	25	61	BZ
DeKhotinsky cement ((hard)	25	106	BZ
Carnauba wax		25	107	BZ
Shellac		25	107	BZ
"Opal wax 20"		25	119	BZ
"Night Blue" 36%	of monolayer		35 °	VV
Plates coated with ole	ic acid. see			L. ICT
Plates coated with var	ious (65) substances.	see		N. ICT
Carbon tetrachloride ((liquid), see			CA, ICT
Glyptol resin ("Glyptol DeKhotinsky cement (Carnauba wax	(hard) of monolayer ic acid, see ious (65) substances.	25 25 25 25	106 107 107 119	BZ BZ BZ BZ VV L, ICT N, ICT

a References:

Ablett, A., Phil. Mag. (6), 46, 244-256 (1923).

BH

BM

Ablett, A., Phil. Mag. (6), 40, 244-250 (1925).
Bosanquet, C. H., and Hartley, H., Idem, 42, 456-461 (1921).
Bartell, F. E., and Miller, M. A., J. Phys'l Chem., 40, 889-894 (1936).
Bartell, F. E., and Zuidema, H. H., J. Am. Chem. Soc., 58, 1449-1454 (1936).
Coghill, W. H., and Anderson, C. O., U. S. Bur. Mines Tech. Paper No. 262, 1923.
DeWitt, C. C., J. Am. Chem. Soc., 57, 775-776 (L) (1935).
Young, T. F., and Harkins, W. D., Int. Crit. Tables, 4, 434 (1928).
Kneen, E., and Benton, W. W., J. Phys'l Chem., 41, 1195-1203 (1937). BZ CA DW ICT

KB

L

Kneen, E., and Benton, W. W., J. Physis Chem., 41, 1175-1203 (1957).

Langmuir, I., Trans. Faraday Soc., 15, Pt. 3, 62-74 (1920).

Mack, G. L., J. Phys'l Chem., 40, 159-167 (1936).

Nietz, A. H., Idem, 32, 255-269 (1928).

Talmud, D., and Lubman, N. M., Z. Physik. Chem. (A), 148, 227-232 (1930).

Voet, A., and Van Elteren, J. F., Rec. Trav. Chim. Pays-Bas, 56, 923-926 (1937).

^b If copper sulfide is ground under water, $\theta = 0$; but if the same ground surface is exposed to air. θ becomes finite.

For a drop of water on a glass plate on which is an adsorbed layer of Night Blue (purest Nachtblau from Dr. Gruebber, Leipzig), $\theta = 35^{\circ}$ if the adsorbed layer \ge 36 per cent of a monolayer. On decreasing the coverage below 36 per cent, θ decreases sharply but continuously, becoming 0 (complete wetting of the plate) when the coverage is 32 per cent of a monolayer.

⁵⁴ Bartell, F. E., Culbertson, J. L., and Miller, M. A., J. Phys'l Chem., 40, 881-888 (1936).

⁵⁵ Bartell, F. E., and Miller, M. A., J. Phys'l Chem., 40, 889-894, 895-904 (1936).

Effect of Overlying Gas upon the Surface-tension.

The conclusions of B. Tamamushi ⁵⁶ regarding the effect of the overlying gas upon the tension of the gas-liquid surface appear to be in direct conflict with those of A. Ferguson, ⁵⁷ but the conflict may be less serious than it appears.

Ferguson used Jaeger's method, based upon the pressure required to blow and detach a bubble, and compared the tension when the gas is air with that when it is CO₂. In each case the pressure of the gas was very slightly greater than 1 atm. He "found that whether the liquids were gasfree or partially or completely saturated with either or both of the gases employed, the result was always the same, viz.: the difference between the surface-tensions liquid-air and liquid-CO₂ remained the same, but the absolute values of the surface tensions increased slightly as the liquid became more and more saturated, finally reaching a steady value." ⁵⁷, p. 407

Tamamushi used the method based upon the rise of the liquid in capillary tubes, and studied the effects of several gases upon the surface-tension of each of several liquids. His observations show that the tension of the surface separating a gas-saturated liquid from the gas itself, saturated with the vapor of the liquid, is in every case less than that of the surface separating the pure liquid from its pure vapor. Furthermore, he states that his data, all of which are for pressures not far from 1 atm, satisfy quite closely the empirical relation $(\gamma_v - \gamma_g)/\gamma_v = KC^{1/3} + b$, where γ_v is the surface-tension of the pure liquid in contact with its pure vapor, γ_g is that of the gas-saturated liquid in contact with the moist gas, C is the mass of gas dissolved in 100 units (mass) of the liquid, and K and b are constants depending upon the liquid, but not upon either the nature or the density of the overlying gas. His data indicate that for water K = 0.0175, b = -0.00052. Actually, the relation is not strictly linear, but the numerical values of K and b decrease as C becomes smaller.

According to Tamamushi's formula, γ_{σ} decreases as C increases, whereas Ferguson states that it increases as the liquid becomes more nearly saturated with the gas. This discrepancy should be investigated with great care. It seems possible that it arises in this manner: Tamamushi did not study the variation of γ_{σ} with the pressure of the gas, which in each case for water lay between 0.73 and 1.0 atm. Hence, his C's are, to a first approximation, proportional to the solubilities of the several gases. In fact, his data for water are more accurately represented by the relation $(\gamma_{\sigma} - \gamma_{\sigma})/\gamma_{\sigma} = 1.54S^{1/3}$, where S = mass of gas dissolved in 100 units (mass) of liquid when the pressure of the gas is 1 atm, than they are by the one he gives. This relation does not conflict with Ferguson's observations. It states that, when the pressure does not vary greatly, the relative depression of the tension is proportional to the cube root of the solubility of the gas. Ferguson's statement is to the effect that for a given

⁵⁶ Tamamushi, B., Bull. Chem. Soc. Japan, 1, 173-177 (1926).

⁵⁷ Ferguson, A., Phil. Mag. (6), 28, 403-412 (1914).

gas and liquid the depression decreases slightly as the amount of gas dissolved in the liquid increases. These two statements are not contradictory, but supplementary.

J. L. R. Morgan and C. E. Davis, 58 working exclusively with an airwater surface, found that saturating the water with air increased the tension of the surface. This agrees with Ferguson's observations. At 0 °C, the increase in going from an unstated initial condition to saturation is given as 0.16 per cent. 58, p. 557, ftn. 2

In the absence of numerical data for the variation of γ_a with the pressure of the gas, it is impossible to reduce the available data to the basis of a constant pressure. Both Ferguson's and Richards and Carver's data obviously refer to a pressure that is very nearly 1 atm; Stocker's data seem to refer to a much lower pressure; the pressures given in Table 229 for Tamamushi's data have been computed from the values he gives for C.

Table 229.—Effect of Overlying Gas upon the Surface-tension of Water

Adapted, with additions, from the compilation of T. F. Young and W. D. Harkins. 59

Remarks in the text should be considered. $\gamma_v = \text{tension of the surface}$ separating the pure liquid from its pure vapor; γ_g = that of the surface separating the liquid saturated with the indicated gas from the gas saturated with the vapor of the liquid. When the gas is air, γ_a is written for γ_g . $\Delta_v \equiv (\gamma_v - \gamma_g)/\gamma_v$, $\Delta_a = (\gamma_a - \gamma_g)/\gamma_a$; that is, $\gamma_g = \gamma_v(1 - \Delta_v) = \gamma_a$ $(1 - \Delta_a)$. p = pressure of the moist gas.

	Unit of $\Delta = 1$	per cent; of	p = 1 atm. Temp	$0. = t ^{\circ}C$	
Gas	t	t	Δ_v	$\Delta_{\mathbf{a}}$	Ref.a
Air	1	20	0.22		T
Air	1	20	0.027 b		RC
CO ₂	1	15		1.1	F.
CO2	0.8	18	0.83	0.61°	\mathbf{T}
CO2	low	20		1.0	S
N_2O	0.9	25.2	0.75 8		Т
H ₂ S	0.7	15.2	1.19		T
H,	low	20		0.0	S
	organic gases				K

[&]quot; References:

Ferguson, A.⁵⁷

Körán, V., Rec. trav. chim. Pays.-Bas, 44, 466-475 (1925).

Richards, T. W., and Carver, E. K., J. Am. Chem. Soc., 43, 827-847 (1921).

Stocker, H., Z. physik. Chem., 94, 149-180 (1920).

Tamamushi, B.⁵⁸ F K

This RC value of Δ_v for air is based upon their reported values, $\gamma_v = 72.75$ and This RC value of Δ_{τ} for all is based upon their reported values, $\gamma_{\tau} = 72.73$ and $\gamma_{\sigma} = 72.73$; and the values of Δ_{τ} for N₈O and H₂S are those reported by T for the respective concentrations of 0.145 and 0.334 g of gas per 100 g of water. From these concentrations and the solubilities of the gases (1.17 and 4.52 g per kg of water per atm) the accompanying values of p have been derived. From the same reports (RC and T), Young and Harkins derived the following values of Δ_{τ} for p = 1 atm: RC, air 0.03_{\circ} ; T, N₈O 0.85; T, H₂S 1.29.
Computed from $\Delta_{\tau} = 0.22$ for air, and 0.83 for CO₂.

²⁰ Morgan, J. L. R., and Davis, C. E., J. Am. Chem. Soc., 38, 555-568 (1916). ²⁰ Young, T. F., and Harkins, W. D., Int. Crit. Tables, 4, 474 (1928).

Miscellanea.—For volume of the water meniscus, see Tables 286 and 287.

Floating bubbles and drops.—Bubbles and drops of water are frequently observed floating upon an air-water surface. In the case of bubbles, the wall of the bubble must be of such a nature that its tension increases as its thickness decreases, which seems to demand that it be a compound film, that it is not pure water. On the case of drops, a blanket of the surrounding medium (air in the case here considered), separating the drop from the surface on which it floats, seems necessary; see L. D. Mahajan, D. B. Seth, C. Anand, and L. D. Mahajan, M. Katalinić, W. and A. R. Hughes, A. Reynolds, T. H. Hazlehurst and H. A. Neville. As is well known an object that is not wetted by water may float on its surface, although much denser than water. The surface is depressed by it and the vertically upward component of the surface-tension supports it. Such seems to be the explanation of the floating of mercury droplets reported by N. K. Adam.

Depression under reduced pressure.—Years ago, K. W. v. Nägeli 67 observed that when there is placed under the receiver of an air-pump a vessel of water into which dips a vertical capillary tube, and the receiver is exhausted, then, under suitable conditions, the meniscus in the capillary is depressed. He showed that this depression is in large part due to the excess of the existing vapor pressure over the meniscus above that over the surface in the large vessel, this excess being due to the resistance encountered by the vapor in streaming through the tube. In many cases, this seemed sufficient to account for the observations. But under certain conditions, especially when the pump was worked rapidly, the depression quite significantly exceeded all that could be accounted for by such difference in the vapor pressure. He also observed that the meniscus in the capillary descended at the same rate as that in a second capillary of the same size, but closed below, and filled to the same distance from the top. This, in connection with the well-known tendency for a stationary meniscus to become stuck to the tube, led him to suggest that as the boundaries of a liquid are approached the molecules assume an arrangement that is more orderly and less mobile than that in the interior. The liquid is thus enclosed in a relatively immobile sheath many molecules thick. The adherence of the sides of the sheath to the walls of the capillary anchors the cap

⁶⁰ Cf. Lord Rayleigh's remarks in the article, "Capillary Action," in "Encyclopedia Britannica," 11th ed., vol. 5, p. 267, 1910.

⁶¹ Mahajan, L. D., Z. Physik, **90**, 663-666 (1934); **84**, 676 (1933); **81**, 605-610 (1933); **79**, 389-393 (1932); Koll. Z., **66**, 22-23 (1934); **69**, 16-21 (1934); Nature, **126**, 761 (1930); **127**, 20 (erratum) (1931); Phil. Mag. (7), **10**, 383-386 (1930).

Seth, J. B., Anand, C., and Mahajan, L. D., Phil. Mag. (7), 7, 247-253 (1929).

⁶⁸ Katalinić, M., Z. Physik, 38, 511-512 (1926); Nature, 127, 627-628 (1931).

⁶⁴ Hughes, W. and A. R., Nature, 129, 59 (1932).

^{84a} Reynolds, O., "Papers on Mechanical and Physical Subjects," vol. 1, 413-414, Cambridge Univ. Press, 1900 ← Proc. Manchester Lit. Phil. Soc., 21, 1-2 (1882).

⁶⁵ Hazlehurst, T. H., and Neville, H. A., J. Phys'l Chem., 41, 1205-1214 (1937).

⁶⁶ Adam, N. K., Nature, 123, 413 (1929).

er v. Nägeli, K. W., Sitz-Ber. Bayer. Akad. Wiss. München, 1866 I, 353-376, 473-492, 597-627 (1866).

forming the meniscus, and as the outer molecules evaporate from the cap a corresponding number is added to its interior face, and thus the cap may be slowly depressed down the tube, perhaps to a point below any that can be accounted for by the vapor pressure alone. The original articles should be studied, and the subject reinvestigated.

Transition layers.—W. D. Harkins and H. M. McLaughlin ⁶⁸ have published values for the thickness of the hypothetical film of pure water that forms the air-liquid surface of an aqueous solution of NaCl. The values given vary from 4.0 to 2.3A, depending upon the concentration (1A = 10-8 cm). But F. Lark-Horovitz and J. E. Ferguson ⁶⁹ have reported observations that indicate that the surface layer of such a solution is not pure water, but a solution much more dilute than the body of the liquid.

From a study of the reflection of polarized light by a water surface, C. V. Raman and L. A. Ramdas⁷⁰ concluded that the transition layer between water in bulk and its overlying vapor is 5.3A. They quote the late Lord Rayleigh as having inferred from similar observations a thickness of 3.0A. Similar observations by J. H. Frazer ⁷¹ on the reflection from a glass surface covered with various amounts of adsorbed water indicate that the transition from a glass surface to a water surface is complete when the thickness of the adsorbed water is 3A.

Surface films.—Interesting summaries of the properties of foreign films upon the surface of water have been published by A. Marcelin,⁷² and by H. E. Devaux.⁷³

Certain data for films of water adsorbed on solids, and estimates of the thickness of the layer of water that may be modified by the action of an adjacent solid, have been considered already (p. 513). For additional observations bearing on the subject, see B. Derjaguin,⁷⁴ T. Ihmori,⁷⁵ J. M. Macaulay,⁷⁶ S. Procopiu,⁷⁷ B. H. Wilsdon, D. G. R. Bonnell, and M. E. Nottage.⁷⁸

Relations between the surface-tension and other properties.—Among the various suggested relations between the surface-tension and other properties of a liquid may be mentioned the empirical ones announced by P. Walden,⁷⁰ by R. K. Sharma,⁸⁰ and by D. Silverman and W. E. Roseveare,⁸¹ and the theoretical one by S. C. Bradford.⁸²

- 68 Harkins, W. D., and McLaughlin, H. M., J. Am. Chem. Soc., 47, 2083-2089 (1925).
- ⁶⁰ Lark-Horovitz, F., and Ferguson, J. E., Phys. Rev. (2), 42, 907 (A) (1932).
- 70 Raman, C. V., and Ramdas, L. A., Phil. Mag. (7), 3, 220-223 (1927).
- 71 Frazer, J. H., Phys. Rev. (2), 33, 97-104 (1929).
- ⁷² Marcelin, A., "Solutions superficielles, fluides à deux dimensions, et stratifications monomolécaires," 163 pp., 86 Figs., bibliog. of 90 entries. Presses Univ. de France, Paris, 1931.
 - 78 Devaux, H. E., Jour. de Phys. (7), 2, 237-272 (1931). Bibliog. of 95 entries.
 - 74 Derjaguin, B., Nature, 138, 330-331 (L) (1936).
 - 75 Ihmori, T., Ann. d. Physik (Wied.), 31, 1006-1014 (1887).
 - 76 Macaulay, J. M., Nature, 138, 587 (L) (1936).
 - 77 Procopiu, S., Compt. rend., 202, 1371-1373 (1936).
- ⁷⁸ Wilsdon, B. H., Bonnell, D. G. R., and Nottage, M. E., Trans. Faraday Soc., 31, 1304-1312 (1935); 32, 570 (1936); Nature, 135, 186-187 (L) (1935).
- ⁷⁰ Walden, P., Z. physik. Chem., 65, 129-225, 257-288 (1908); 66, 385-444 (1909); Z. Elektrochem., 14, 713-724 (1908).
- 80 Sharma, R. K., Chem. Abs., 20, 2267 (1926) ← Quart. Jour. Indian Chem. Soc., 2, 310-311 (1925).

The most important, however, is that which S. Sugden 83 called the "parachor" and denoted by P. He defines it by means of the formula $P = \gamma^{1/4} M/(D-d)$, where $\gamma = \text{surface-tension of the liquid-vapor sur-}$ face, M = molecular weight of the vapor, and D and d are the densities of the liquid and vapor, respectively. For many substances, P is almost independent of the temperature, and its value can be obtained by summing certain constants, each characteristic of a chemical element or of a type of structure that enters into the make-up of the molecule. Sugden has stated that within 3 per cent $P = 0.78 V_o$, where V_o is the critical volume of a gram-mole. See also, N. K. Adam.84

Movements of bubbles.—Although the phenomena accompanying the motion of bubbles in a liquid depend in part upon the surface tension of the liquid-gas boundary, their discussion is scarcely pertinent to the present compilation. Several papers treating of them have, however, happened to come to the compiler's attention.85

Voltaic effects.—Voltaic effects suggesting that the capillary layer or, more exactly, the portion of the liquid that is elevated in a tube, as a result of capillary action, differs voltaically from the liquid in bulk have been reported by E. Torporescu.86

Stability of doubly gas-faced liquid films.—A clean, uncoated film of a pure liquid, each face in contact with a gas, is incapable of static equilibrium. A dynamic equilibrium, resulting from localized evaporation and streaming, is however possible, and has been recently considered by H. A. Neville and T. H. Hazlehurst, Ir.87

85. Solubility of Selected Gases in Water

Definitions and Symbols.

By the *solubility* of substance A in substance B is meant the amount of A that must be added to a unit amount of B in order to produce a solution that will be in equilibrium with an excess of A under the existing conditions. Such a solution is said to be saturated with A under those conditions. The definition of solubility is not concerned with the state of A after solution has occurred, although that will, in general, affect the magnitude of the solubility, and may perhaps give rise to special effects.

⁸¹ Silverman, D., and Roseveare, W. E., J. Am. Chem. Soc., 54, 4460 (1932).

⁸² Bradford, S. C., Phil. Mag. (6), 48, 936-947 (1924).

Sugden, S., J. Chem. Soc. (London), 125, 1177-1189 (1924).
 Adam, N. K., "The Physics and Chemistry of Surfaces," Oxford Univ. Press, 1930. **Solution Research Commission of Surfaces, Oxfold Univ. Press, 1930.

**Solution of Surfaces, Oxfold Univ.

⁸⁶ Torporescu, E., Bull. Math. et Phys., Bucarest, 6, 40-41 (1936); Compt. rend., 202, 1672-1674 (1936).

⁸⁷ Neville, H. A., and Hazlehurst, T. H., Jr., J. Phys'l Chem., 41, 545-551 (1937).

⁸⁸ Int. Crit. Tables, 3, 254-255 (1928).

⁵⁰ Loomis, A. G., Int. Crit. Tables, 3, 255-261 (1928).

2. By the coefficient of absorption of a gas in a liquid is meant the rate at which the solubility of the gas in it increases with the partial pressure of the gas in the overlying gaseous phase, the temperature remaining

Table 230.—Mean Coefficients of Absorption (0 to P) of Selected Gases by Water

With but few exceptions the values in this table have been derived from the mean molecular coefficients given in Table 232. The two are connected through the relations

 $r = xM_g/(1-x)M_l$ and $f = xM_g/(M_l - (M_l - M_g)x)$ where M_{g} and M_{l} are the formula-weights of the gas and of $H_{2}O$, respectively, x = mole-fraction of the gas in the solution, r and f = mass of gas dissolved in unit mass of water, and in unit mass of the solution, respectively (see p. 535+). The difference in the units of pressure used in the two tables must be considered. For all the gases in this table, excepting CO₂ and NH₃, x is negligibly small in comparison with unity, and consequently r=f.

Unless otherwise indicated, the values here given apply when the partial pressure (P) of the gas is 1 atm; and, excepting NH₃, it is probable that r/P and f/P are independent of P if that does not much exceed 1 atm. But no data are available for the radioactive gases except at very low pressures. For variation of the coefficients with the pressure, see Tables 233 and 234.

Units: Of r/P [f/P] = 1 mg of gas per kg of water [of solution] per atm; of $\lambda = 1$ cm³ of gas under existing conditions per cm³ of solution. Temperature = t °C; $M_i = 18.0154$.

I. Noble gases.

An Actinon (actinium emanation). At exceedingly low partial pressures $\lambda = 2.88$

Rn Radon (radium emanation). See end of Section II.
Th Thoron (thorium emanation). At exceedingly low partial pressures, $\lambda = 1.80$ See also A. Klaus. 4

$M_g \rightarrow t$		Argon		= f/P	Helium 4.00	
0 5	94.1	103.0 90.46	.,.	(1.69) 1.647	1.725	,
10 15	74.8	80.60 73.07	66.1	1.606 1.571	1.768	1.59
20	62.3	67.04	59.9	1.539	1.777	1.57
25 30 35 40	53.5	62.18 58 15 54.58	56.0 53.3	1.504 1.476	1.791	1.55 1.54
40 45 50	48.5	51.57 48.84 46.35			1.836 1.932	
Ref. ₁ s Ref. ₂ s	W97, W06 V27	E V27	I.	CER V27	A ^b V27	L
a₀° a₁° a₂°		-110 +1.38 -9.7			$-110 \\ +1.38 \\ -10.5$	

⁹⁰ Metschl, J., J. Phys'l Chem., 28, 417-437 (1924).

⁹¹ Manchot, W., Z. anorg. allgem. Chem., 141, 38-4' (1924).

Table 230.—(Continued)

$Gas \rightarrow M_g \rightarrow t$	Krypton 82.9	Neon 20.2	= f/P	Xenon 130.2
0 5 10	370.0	9.8	*/-	1406 1186
10 15	295.5	10.8	9.7	1005 862
20	231.5	26.1	9.4	741
25 30 40 45.45	188.8 160.2	49.1 95.2	9.1 8.9	570 467 424
50 Ref.1ª Ref.2ª	141.6 ^d A V27	89.7 A ^b V25, V27	L	A V27
$a_0^{\ \sigma} \ a_1^{\ \sigma} \ a_2^{\ \sigma}$	-108 +1.35 -9.4	$-110 \\ +1.38 \\ -9.9$		-103 + 1.30 - 9.0

II. Simple gases.

(For noble gases, except radon, see Section I.)

1	2	3	4	5	6	7
$Gas \rightarrow M_g \rightarrow$	$H_2 H_3$	ydrogen)154	Argon-f	ree N ₂ e 16	"Atmosp	heric'' N ₂ 1 .016
ŧ			r/P = 1	F/P		
0	1.936	1.931 *	28.99	28.90	29.50	29.43
1	1.919	1.912			28.81	28.72
2	1.902	1.893			28.14	28.03
1 2 3 4	1.885	1.874			27.49	27.34
	1.870	1.856			26.89	26.69
5 6 7 8 9	1.853	1.839	25.85	25.64	26.29	26.06
6	1.838	1.821			25.73	25.47
7	1.823	1.805			25.18	24.87
8	1.808	1.789			24.65	24.31
	1.793	1.773			24.16	23.78
10	1.780	1.759	23.29	22.87	23.70	23.27
11	1.765	1.745			23.24	22.80
12	1.751	1.732			22.82	22.34
13	1.737	1.718			22.41	21.89
14	1.725	1.706			22.03	21.50
15	1.712	1.694°	21.30	20.72	21.66	21.08
16	1.701	1.682			21.30	20.70
17	1.688	1.672			20.96	20.34
18	1.676	1.663			20.62	19.99
19	1.665	1.649			20.32	19.66
20	1.654	1.638 ". "	1.638°	19.01 "	20.01	19.35
21	1.644	1.626			19.73	19.05
22	1.634	1.616			19.45	18.77
23	1.624	1.604			19.19	18.48
24	1.614	1.592			18.93	18.22
25	1.605	1.582 0. 0	1.582 °	17.68"	18.67	17.98
26	1.597	1.572			18.44	17.72
27		1.562			18.20	17.49
28 29		1.552			17.98	17.27
		1.543			17.76	17.05
30		1.535	17.26	16.53	17.56	16.83
35		1.508	15.97	15.52	16.60	15.79

Table 230.—(Continued)

1	2	3	4	5	6	7
$Gas \rightarrow t$	H, 1	Hydrogen	Argo:	P = f/P	"Atmos	pheric' N ₂ /
40 45 50	·	1.490 1.475 1.463	15.53 14.79 14.12	14.68 14.02 13.55	15.77 15.01 14.34	14.92 14.26 13.79
60 70 80 90 100		1.463 1.47 1.48 1.49 1.50				13.01 12.49 12.32 12.30 12.26
Ref1ª Ref2ª	Ti	W91a, W92b W92a			Fox W	91b, W92b W92a
$M_{g} \rightarrow 0$		O ₂ Oxygen ^h ————————————————————————————————————		$O_3 Ozonef$ $= f/P$		Radon / ———————————————————————————————————
0		69.82 67.94 66.15° 64.42 62.78	70.29 68.37 66.55 64.82 63.19	1373.	5049.	5130. 4920. 4710. 4530. 4360.
2 3 4 5 6 7 8 9		61.21 59.68 58.26 56.88 55.57	61.64 60.17 58.76 57.41 56.12	1220.	4080.	4190. 4020. 3870. 3720. 3580.
10 11 12 13 14		54.30 53.11 51.96 50.85 49.81	54.87 53.69 52.55 51.47 50.44	1075.	3339.	3450. 3320. 3180. 3070. 2960.
15 16 17		48.80 47.85 46.93	49.43 48.47 47.55	924.		2860. 2750. 2650.
18 19		46.04 45.20	46.67 45.80	974.*		2560. 2470.
20 21 22 23 24	44.36 43.55 42.76 42.05 41.24	44.68° 43.62 42.86 42.12 41.42	44.98 44.20 43.45 42.72 42.02	709.	2360.	2380. 2300. 2240. 2160. 2100.
25 26 27 28 29	40.53 39.87 39.20 38.57 37.99	40.73 ° 40.08 39.41 38.78 38.13	41.35 40.70 40.07 39.47 38.89	583.		2040. 1980. 1920. 1850. 1830.
30 35	37.40 35.05	37.50	38.33 35.80	445.3 323.6	1780.	1780. 1580.
40 45 50	33.18 31.53 30.20		33.67 31.88 30.37	222.3 149.1 97.1 ¹	1410. 1180.	1420. 1300. 1200.
60 70 80 90 100	28.26 26.77 25.84 25.42 25.31		30.0.	0	1050. 960. 880. 840. 800.	1030. 920. 860. 830. 820.**
Ref ₁ ^a Ref ₂ ^a	W91b W06	W89, W92b W92a, W88	Fox	Mt	StM	Sz.

Table 230.—(Continued)

III. Gaseous compounds.

In all cases the partial pressure (P) of the gas either was very nearly 1 atm or lay in a region, including 1 atm, throughout which r/P and f/P are independent of P. Excepting NH₃, $(\partial r/\partial P)_t$ and, consequently, $(\partial f/\partial P)_t$ and $(\partial x/\partial P)_t$, are constant from a very low pressure to a value of P that is well above 1 atm.

$\begin{array}{c} 1 \\ Gas \rightarrow \\ M_g \rightarrow \\ t \end{array}$	2 COn 28.000 r/P	44.	4 0000 /P	5	6 CO₂º 4.000 f/P		8 H ₇ p 031 10-0f/P
0 1 2 3 4	44.19 43.17 42.17 41.19 40.25	3364. 3232. 3109. 2998. 2892.	ľ	3352 3221 3100 2989 2884	y/r	0.895 0.872 0.846 0.820 0.799	0.472 0.464 0.457 0.450 0.443
5 6 7 8 9	39.34 38.45 37.60 36.76 35.97	2789. 2696. 2605. 2518. 2431.		2781 2689 2599 2511 2425		0.778 0.759 0.741 0.720 0.703	0.438 0.431 0.425 0.419 0.412
10 11 12 13 14	35.19 34.46 33.76 33.07 32.42	2346. 2267. 2198. 2127. 2060.		2342 2262 2192 2123 2056		0.685 0.668 0.652 0.636 0.620	0.407 0.401 0.394 0.388 0.383
15 16 17 18 19	31.80 31.20 30.62 30.05 29.56	1999. 1976. 1880. 1824. 1770.		1994 1935 1877 1821 1768		0.604 0.587 0.572 0.560 0.548	0.376 0.369 0.364 0.359 0.353
20 21 22 23 24	20.02° 28.55 28.22 27.65 27.24	1722. 1672. 1623. 1579. 1536.	1648. 1582. 1512.	1718 1669 1621 1576 1534	1645 1580 1509	0.534 0.522 0.510 0.500 0.488	0.348 0.342 0.337 0.333 0.328
25 26 27 28 29	26.84° 26.45 26.08 25.73 25.40	1493." 1452. 1419. 1384. 1346.	1439. 1367.	1492 1451 1417 1382 1345	1438 1366	0.479 0.465 0.456 0.448	0.324 0.320 0.313 0.309
30 32 34 35 40	25.08 23.59 22.35	1317. 1170. 1049.	1294. 1224. 1151.	1312 1168 1048	1293 1222 1151		
45 50 60 70 80	21.32 20.42 18.91 18.40 18.38	951. 862. 717.		949 862 716			
90 100 Ref.1 ^a Ref.2 ^a	18.4 18.4 W01 W92a	Bohr Bu	Ku Bu	Bohr Bu	Ku Bu	-	

Table 230.—(Continued)

References: See end of this Section.

The values in this column increase with t, suggesting an error in the observations.

^e Coefficients in the formula $M_{\theta}^{-0.8}\log_{10}\lambda = (273a_1 - a_0)/T + 2.3a_1\log_{10}T + a_2$, where T °K is the absolute temperature. For krypton at 60 °C, r/P = 132.2.

This is argon-free atmospheric nitrogen (see next note). The values in columns 4 and 5 have been respectively computed from those of columns 6 and 7.00 Apparently the only corresponding data available in 1927 for chemically prepared nitrogen were those reported by Braun (1900), Just (1901), and Adeney and Becker (1919); see Coste, who seems to have overlooked Just's work. None since that date have come to the attention of the compiler. Of these, Loomis included only Just's, which are here given in footnote g. Some of the values by the others are here given, subscripts indicating the gas; they do not accord well with those given in the main table.

Braun									
t	0	15	20	25	t	2.5	3.5	20	25
$(r/P)_{\rm H}$	2.130	1.854	1.715	1.576	$(r/P)_0$	63.29		43.46	40.51
$(r/P)_{N}$	27.22	22.41	20.31	17.94	$(r/P)_N$		27.82	19.90	18.78

Atmospheric nitrogen is the residue of air from which O2, CO2, NH3, and H2O

have been removed.

"Just's data (J) yield the following values of r/P, all included in Loomis's compilation (ICT). The nitrogen was prepared chemically, not from the atmosphere.

t	H_2	N_2	CO	CO_2
20	1.67₅	19.90	30.18	
25	1.64	18.77	27.59	1488

And Findlay et al. (F) give for CO₂ at 25 °C the value 1480 for the range 270 to

1350 mm-Hg.

The data in column 2 were obtained by measuring the volume of O₂ absorbed by a given volume of water; those in 3, by titrating the O₂ contained in air-saturated

'The data for ozone refer probably to a partial pressure of about 50 mm-Hg. By

an indirect computation, Rothmund (Ro) found 1057 at 0 °C.

The data for radon refer to an exceedingly low partial pressure, of the order of on the observations of Boyle, H, Ko, Mache, Ra, and Tr. The coefficients of Valentiner's formula (see note c) for Rn are $a_0 = -95$, $a_1 = 1.20$, $a_2 = -8.25$.

This value for O_a is from (FT).

'At 55 °C, r/P = 56.0 for O₃.

^m This value for Rn is for t = 97 °C.

" For CO, r/P = f/P, essentially.

The values in columns 3 and 4, respectively, represent the same data as those

in 5 and 6.

The values in columns 7 and 8 represent the same data and apply solely to P=1 atm. The ICT formulas by means of which they were computed ($\log_{10}K=0.05223A/T+B$, $A=-937_0$ and $B=4.98_7$ from 0 to 10 °C, and $A=-1074_0$ and $B=5.23_8$ from 14 to 28 °C) seem to have been derived from the observations of G. Calingaert and F. E. Huggins, Jr., E. Klarmann, F. M. Raoult, on A. Smits and S. Postma.101

⁹² Findlay, A., and associates, J. Chem. Soc. (London), 97, 536-561 (1910); 101, 1459-1468 (1912); 103, 636-645 (1913); 107, 282-284 (1915).

ee v. Hevesy, G., Physik. Z., 12, 1214-1224 (1911)-J. Phys'l Chem., 16, 429-450 (1912).

P4 Klaus, A., Physik. Z., 6, 820-825 (1905).

BE Valentiner, S., Z. Physik, 42, 253-264 (1927).

⁹⁰ See Loomis, Int. Crit. Tables, 3, 256 (1928).

⁹⁷ Boyle, R. W., Phil. Mag. (6), 22, 840-854 (1911).

⁸⁸ Calingaert, G., and Huggins, F. E., Jr., J. Am. Chem. Soc., 45, 915-920 (1923).

[™] Klarmann, E., Z. anorg. allgem. Chem., 132, 289-300 (1924).

constant. This quantity does not appear in the tabulations except incidentally where it happens to coincide with the mean coefficient. That frequently occurs.

3. By the mean coefficient of absorption of a gas in a liquid when the partial pressure of the gas is increased from p_1 to p_2 is meant $(S_2 - S_1)/(p_2 - p_1)$, where S_1 and S_2 are the solubilities at p_1 and p_2 , respectively, the temperature being the same in each case. The quantity commonly tabulated and called (unfortunately) the coefficient of absorption is the

Table 231.—Solubility of Air in Water

Adapted from the compilation by A. G. Loomis. 102

(See also, Table 232, part III. For solubility of air in sea-water, see Table 235.)

By "air" is meant atmospheric air that has been freed from CO_2 and NH_3 . r = mass of gas that is contained in a unit mass of water when in equilibrium with air at a pressure of 1 atm and at the indicated temperature.

 v_0 = the volume under standard conditions (0 °C and 1 atm) of the gas that is contained in a unit volume of water when in equilibrium with air at a pressure of 1 atm and at the indicated temperature; $v_0 = r_{Pl}/\rho_0$.

 Σ = total volume of gas (0 °C and 1 atm) = sum of the corresponding values in the next two preceding columns.

The molecular weight of air has been taken as 28.96; the density of O_2 under standard conditions, as 1.4290_4 g per liter, and that of "atmospheric" N_2 (including the inert gases), as 1.2568 g per liter.¹⁰³

Unit of r = 1 mg gas per kg water; of $v_0 = 1$ ml (0 °C, 760 mm-Hg) of gas per liter of water;

Ref.₁ª→	w	01	pressure = 1 atm. Temp.			W04			
Gas→	Airb	Aire	O_2	N2, A, etc.	Σ	O_2	N2, A, etc.	Σ	100 O.
t			r				v ₀	(Σ v_0
0	37.27	37.95	14.56	23.87	38.43	10.19	18.99	29.18	34.91
1	36.34	36.99	14.16	23.26	37.42	9.91	18.51	28.42	34.87
2 3 4	35.40	36.02	13.78	22.68	36.46	9.64	18.05	27.69	34.82
3	34.53	35.14	13.42	22.12	35.54	9.39	17.60	26.99	34.78
	33.71	34.28	13.06	21.59	34.65	9.14	17.18	26.32	34.74
5	32.91	33.44	12.73	21.08	33.81	8.91	16.77	25.68	34.69
6 7	32.12	32.68	12.40	20.59	32.99	8.68	16.38	25.06	34.65
7	31.37	31.93	12.10	20.11	32.21	8.47	16.00	24.47	34.60
8	30.66	31.22	11.80	19 66	31.46	8.26	15.64	23.90	34.56
9	29.97	30.53	11.52	19.23	30.75	8.06	15.30	23.36	34.52
10	29.32	29.88	11.25	18.82	30.07	7.87	14.97	22.84	34.47
11	28.71	29.26	10.99	18.42	29.41	7.69	14.65	22.34	34.43
12	28.11	28.66	10. 7 5	18.05	28.80	7.52	14.35	21.87	34.38
13	27.53	28.10	10.51	17.68	28.19	7.35	14.06	21.41	34.34
14	<i>27.</i> 00	27.54	10.28	17.33	27.61	7.19	13.78	20.97	34.30
15	26.49	27.03	10.07	17.00	27.07	7.04	13.51	20.55	34.25
16	25.99	26.53	9.86	16.67	26.53	6.89	13.25	20.14	34.21
17	25.51	26.06	9.66	16.36	26.02	6.75	13.00	19.75	34.17
18	25.07	25.59	9.46	16.07	25.53	6.61	12.77	19.38	34.12
19	24.61	25.16	9.28	15.78	25.06	6.48	12.54	19.02	34.08

¹⁰⁰ Raoult, F. M., Ann. de chim. et phys. (5), 1, 262-274 (1874).

¹⁰¹ Smits, A., and Postma, S., Proc. Akad. Wet. Amsterdam, 17, 182-191 (1914).

¹⁰² Loomis, A. G., Int. Crit. Tables, 3, 255-261 (257-258) (1928).

Table 231.—(Continued)

Ref.₁ª→	wo	1		W04			W04				
Gas→	Airb	Aire	O ₂	N ₂ , A, etc.	Σ	O ₂ 1	N2, A, etc.	Σ (100 O ₂		
20	24.22	24.74	9.11	15.51	24.62	6.36	12.32	18.68	Σ) _{v₀} 34.03		
21	23.81	24.44	8.92	15.24	24.16	6.23	12.11	18.34	33.99		
22	23.42	23.95	8.75	14.99	23.74	6.11	11.90	18.01	33.95		
23	23.06	23.58	8.59	14.73	23.32	6.00	11.69	17.69	33.90		
24	22.71	23.20	8.44	14.48	22.92	5.89	11.49	17.38	33.86		
25	22.34	22.83	8.28	14.24	22.52	5.78	11.30	17.08	33.82		
26 27	22.04 21.72	22.51 22.16	8.1 5 7.98	14.02 13.80	22.15 21.78	5.67 5.56	11.12 10.94	16.79 16.50	33.77 33.73		
28	21.43	21 83	7.83	13.56	21.39	5.46	10.75	16.21	33.68		
29	21.13	21.50	7.69	13.32	21.02	5.36	10.56	15.92	33.64		
30	20.86	21.19	7.55	13.10	20.65	5.26	10.38	15.64	33.60		
35	19.55					0.20					
40	18.48										
45	17.66										
50	17.00										
60	15.98										
70 80	15.31 14.96					•	•				
90	14.86										
100	14.97										

"References: See end of this Section.

 b Calculated from data for O_{2} and N_{2} with correction for constant amount of argon. 104

^o Calculated from the O₂-content of water saturated with air and of the air expelled from the saturated water by heating.

mean coefficient for the range 0 to p; i.e., it is S/p. Throughout the range in which S/p is independent of p, the mean coefficient coincides with the coefficient itself. When S is expressed in terms of the mole-fraction (x) of the gas in solution, we shall call S/p the mean molecular coefficient of absorption over the range 0 to p.

4. By the coefficient of solubility (λ) of a gas in a liquid is meant the volume of the gas, as measured under the conditions existing in the gas phase, that is contained in unit volume of the saturated solution. This terminology essentially agrees with ordinary practice, but in *International Critical Tables* λ is called the Ostwald absorption coefficient, and the definition is so worded as to make it appear as a partition coefficient.

The solubility of a gas may be expressed in several ways, to each of which corresponds a different set of values for the several coefficients. Thus a complex and somewhat confusing terminology has arisen; this condition is aggravated by a lack of unanimity regarding the actual significance of the terms and symbols commonly employed. For this reason the definitions just given will be adhered to, and in the presentation of the data but one (λ) of the symbols commonly used for denoting the coefficients will be employed. The symbols that will be used are as follows:

¹⁰⁸ See Int. Crit. Tables, 3, 3 (1928).

¹⁰⁴ See Fox, C. J. J., Trans. Faraday Soc., 5, 68-87 (1909).

 $r \equiv m_g/m_l$ = ratio of the mass of the dissolved gas to the mass of the pure liquid in which it is dissolved.

 $f \equiv m_g/(m_l + m_g)$ = ratio of the mass of the dissolved gas to the total mass of the solution.

 $x \equiv n_g/(n_l + n_g) = \text{mole-fraction of the gas in the solution} = \text{ratio of}$ the number of gfw of the gas in solution to the sum of that number and the number of gfw of the liquid in which the gas is dissolved. The formula of the gas is to be taken as that pertinent to its pure gaseous state.

 $\lambda = V_g/V_s$ = the volume, under the existing conditions of temperature

and pressure, of the gas contained in unit volume of the solution.

It will be noticed that r, f, x, and λ are all measures of solubility. When the densities of the solution (ρ_{θ}) , of the pure liquid (ρ_{l}) , and of the pure gas (ρ_{θ}) , all under the existing conditions of temperature and pressure, are known, then r, f, x, and λ can be readily interconverted, the formula weights (M_{θ}, M_{l}) of the gas and the liquid being known. If r is negligible as compared with unity, then $r = f = M_{\theta}x/M_{l}$; r is so negligible in every case considered in this section, excepting only CO_{2} and NH_{3} .

 p_g and p_v mm-Hg, or P_g and P_v atm, are the partial pressures, in the gas phase, of the gas and of the vapor of the liquid, respectively; $p_g = 760P_g$, $p_v = 760P_v$.

 $P_t = P_y + P_v = \text{total pressure}.$

 V_0 = volume of the dissolved gas at 0 °C and 1 atm, $V_{0\rho_0} = m_g$; if the gas were ideal and the actual temperature of the system were t °C, then $V_g = V_0(273.1 + t)/(273.1)P_g$.

 V_l = volume of the pure liquid, at the existing temperature and pressure, contained in the solution, $V_{lPl} = m_l$.

 ρ_0 = density of the pure gas at 0 °C and one normal atm.

The relations connecting the symbols used in the International Critical

Table 232.—Mean Molecular Coefficient of Absorption (0 to p) of Selected Gases by Water

Adapted from the compilation by A. G. Loomis. 105

x/p is the reciprocal of the quantity designated by K in Loomis's compilation; $x = n_g/(n_g + n_l)$, where n_g and n_l denote the number of formulamolecules of gas and of H_2O , respectively, contained in a given amount of the solution that is in equilibrium with the gas under the partial pressure p, the rest of the pressure arising from water-vapor with which the gas is saturated. In general, x/p varies with p; but if p does not much exceed 760 mm-Hg it is probable that x/p is independent of p for all the gases in this table except NH_3 . But no data are available for Rn except at very low pressures.

Except as indicated, the values given here apply when p = 760 mm-Hg. Unit of x/p = 1 formula-molecule of gas per 10° formula-molecules of (gas + H₂O) contained in the solution per mm-Hg. Temperature = t°C.

¹⁰⁵ Loomis, A. G., Int Crit. Tables, 3, 255-261 (1928).

Table 232.—(Continued)

I. Noble gases.

Rn, Radon (radium emanation). See end of Section II-Simple gases.

$G_{as o t}^{1}$	A	rgon	4 Heli	uma5	Krypton	7 Neona	8 Xenon
0 5	55.9	61.16 53.73	(10) 9.76	10.22	105.8	11.5	256 216
10 15	44.4	47.87 43.40	9.52 9.31	10.48	85.5	13.0	183 1 <i>5</i> 7
20	37.0	39.82	9.12	10.53	66.2	14.8	135
25 30 35	31.8	36.93 34.54 32.42	8.91 8.75	10.65	54.0	17.5	103.8
40 45	28.8	30.63 29.01		10.88	45.8	22.9	85.0 77.2°
50 60		27.53		11.45	40.5 37.8	34.0	
Ref.1° Ref.2°	W97, 06	V27 E	CEB V	A 27	A V27	A V25, 27	V27

II. Simple gases.

For noble gases, excepting radon, see section I.

$Gas \rightarrow t$	H ₂ Hy	drogen —	$\frac{4}{N_2 \operatorname{Arge}}$	on-freed	6 N ₂ Atm	ospheric*
0 1 2 3 4	22.77 22.57 22.37 22.18 21.99	22.72 22.49 22.27 22.04 21.83	24.53	24.46	24.96 24.37 23.81 23.26 22.75	24.90 24.29 23.70 23.13 22.58
5 6 7 8 9	21.80 21.62 21.44 21.26 21.09	21.63 21.42 21.23 21.04 20.85	21.87	21.70	22.24 21.77 21.30 20.86 20.44	22.05 21.55 21.04 20.57 20.12
10 11 12 13 14	20.93 20.76 20.60 20.44 20.29	20.69 20.53 20.37 20.21 20.07	19.71	19.35	20.05 19.67 19.31 18.96 18.64	19.69 19.29 18.90 18.52 18.19
15 16 17 18 19	20.14 20.01 19.86 19.72 19.58	19.93 19.79 19.66 19.56 19.40	18.02	17.53	18.32 18.02 17.74 17.45 17.19	17.84 17.51 17.21 16.91 16.63
20 21 22 23 24	19.46 [*] 19.34 19.22 19.10 18.99	19.26 ' 19.13 19.00 18.86 18.73	16.66*	16.081	16.93 16.69 16.46 16.23 16.02	16.37 16.12 15.88 15.64 15.42
25 26 27 28 29	18.88 <i>†</i> 18.78	18.61 ' 18.49 18.37 18.26 18.15	15.54 <i>*</i>	14.96 <i>†</i>	15.80 15.52 15.40 15.21 15.03	15.21 14.99 14.80 14.61 14.42
30		18.05	14.61	13.99	14.86	14.24

Table 232.—(Continued)

		Table	232(00)	itinuea)		
1	2	3	4	5	6	7
Gas→ t	H ₂	Hydrogen-		on-freed	N ₂ At:	mospheric•
35		17.74	13.82	13.13	14.04	13.36
40		17.52	13.14	12.42	13.35	12.62
45		17.35	12.52	11.87	12.70	12.07
50 60		17.21 17.21	11.95	11.46	12.13	11.65 11.00
70		17.3				10.57
80		17.4				10.43
90		17.5				10.4
100	TD*	17.7			77. 37.	10.5
Ref. ₂ °	Ti	W91a, W92b W92a			Fox W	791b, W92b W92a
$Gas \rightarrow t$	2	O ₂ Oxygen ⁹	4 	O ₃ Ozone ^h	Rn R	ladon' 7
0	•	51.72	52.07	678.0	539.1	548.
1		50.33	50.64			526.
2		48.99 47.72	49.30 48.02			503. 484.
4		46.51	46.81			466.
5		45.34	45.66	602.4	435.7	447.
6		44.21	44.57			429.
7		43.16	43.52			413.
1 2 3 4 5 6 7 8 9		42.13 41.16	42.52 41.57			397. 382.
10		40.22	40.65	530.8	356.5	368.
11		39.34	39.77	300.0	050.5	354.
12		38.49	38.93			340.
13		37.66 36.96	38.13			328.
14 15		36.86 36.15	37.36 36.62	456.4		316. 305.
16		35.45	35.90	430.4		294.
17		34.77	35.22			283.
18		34.10	34.57	480.8		273.
19 20	22.06	33.48 32.88	33.93	350.1	252.	264. 254.
21	32.86 32.26	32.30 32.31	33.32 32.74	330.1	252.	234. 246.
22	31.68	31.75	32.18			239.
23	31.10	31.20	31.64			231.
24	30.55	30.68	31.12	207.0		224.
25 26	30.03 29.53	30.18 29.69	30.63 30.15	287.8		218. 211.
27	29.04	29.20	29.68			205.
28	28.57	28.73	29.24			200.
29	28.13	28.25	28.81			195.
30 35	27.70	27.78	28.39	219.9 160.8	190.	190. 169.
35 40	25.96 24.58		26.52 24.94	109.8	151.	152.
45	23.36		23.62	47.96*	126.	128.
50	22.37		22.50	73.64		139.
60	20.93			0	112.	110.
70 80	19.83 19.14				102. 94.	98. 92.
90	18.83				90.	89
100	18.75				86.	88.1
Ref.1° Ref.2°	W91b W06	W89, W92b W9 2a	Fox	Mt	StM	Sz

Table 232,-(Continued)

III. Air and gaseous compounds.

In all cases here considered, the partial pressure either was very nearly 1 atm or lay in a region, including 1 atm, throughout which x/p remains essentially constant; excepting NH₈, x/p is independent of p from very low pressures up to a value well above 1 atm.

1 Gas→	Air	3 r ^m	4 CO	CC	6	7 NH ₈ n
<i>t</i> 0	30.51	31.06	x/p 37.41	1810		$\frac{10^{-6}x/p}{6.40}$
	29.75	30.28	36.55	1739		6.31
1 2 3 4	28.98	29.48	35.70	1673		6.21
3	28.28	28.76	34.87	1613		6.11
	27.59	28.06	34.08	1556		6.02
5 6 7 8 9	26.94	27.37	33.31	1501		5.94
6	26.30	26.75	32.55	1451		5.86
7	25.68	26.14	31.83	1402		5.78
8	25.09	25.56	31.12	1355		5.69 5.61
	24.53	24.99	30.45	1308		
10 11	23.99 23.49	24.46 23.95	29.79 29.17	1263 1220		5.53 5.45
12	23.00	23.46	28.58	1183		5.37
13	22.53	23.00	28.00	1145		5.29
14	22.10	22.54	27.45	1109		5.21
15	21.68	22.12	26.92	1076		5.13
16	21.27	21.72	26.41	1044		5.04
17	20.88	21.33	25.92	1012		4.96
18	20.52	20.95	25.44	982		4.90
19	20.14	20.59	25.02	953	007	4.83
20	19.82	20.25	24.57 [†] 24.17	927 900	887	4.75
21 22	19.49 19.17	19.92 19.60	23.89	900 874	852	4.68 4.61
23	18.88	19.30	23.41	850	032	4.55
24	18.59	18.99	23.06	827	814	4.48
25	18.29	18.69	22.72*	804 ^r		4.42
26	18.04	18.42	22.39	782	775	4.34
27	17.78	18.14	22.08	764		4.28
28	17.54	17.87	21.78	745	73 6	4.23
29	17.30	17.60	21.50	725	com	
30	17.07	17.35	21.23	709	697	
35 40	16.00 15.13		19.97 18.92	630 565		
45	14.46		18.05	512		
50	13.91		17.29	464		
60	13.08		16.01	386		
70	12.53		15.58			
80	12.24		15.56			
90	12.2		15.6			
100	12.2		15.6		77	
Ref ₁ ° Ref ₂ °	W01	W01	W01 W92a	Bohr	Ku	

^e In contrast with the others, the values in columns 5 and 7 increase with t, suggesting a serious error in A's data.

^b This xenon value refers to t = 45.45 °C.

References: See end of this section.

⁴ This is argon-free atmospheric nitrogen (see Table 230, notes e and f). Loomis states that the data for columns 4 and 5 have been respectively derived from those of 6 and 7.

Table 232.—(Continued)

 Atmospheric nitrogen is the residue of air from which O2, CO2, NH3, and H2O have been removed.

Just's data (J) yield the following values of x/p, all included in Loomis's compilation. The nitrogen was prepared chemically, not from the atmosphere.

t	H_2	N_2	CO	CO_2
2 0	19.72	16.84	25.55	
25	19.3 ₀	15.88	23.36	802

And Findlay et al. (F) give for CO₂ at 25 °C the value 797, p ranging from 270 to 1350 mm-Hg.

See Table 230, note h.

* Probably for p = 50 mm-Hg. By an indirect computation, Rothmund (Ro) found for ozone x/p = 522.2 at 0 °C.

The pressure for radon was of the order of 0.01 μ -Hg (see Table 230, note j).

This value for On is from FT.
At 55 °C, x/p = 26.67 for On This value for Rn is for 97 °C.

The values in column 2 have been computed from the coefficients of absorption of O2 and of N2, with due attention to the argon content of the atmosphere (see Fox); those in column 3, from the O2 content of water saturated with air, and from the O2 content of the air expelled from the saturated water by boiling.

"These values have been computed by means of the formulas given by Loomis, and apply only to p=760 mm-Hg (see Table 230, note p). For solubility of NH₈ at very low pressure, see remarks under SdW and Wijs in Table 233, note k.

Table 233.—Effect of Pressure on the Solubility of Gases in Water

Adapted, with additions, from the compilation by A. G. Loomis.¹⁰⁵ (See also Table 234.)

 $\lambda = V_g/V_s$ = the volume of gas contained in unit volume of solution in equilibrium with the overlying gas, both volumes being measured under the existing conditions of p and t.

 $r = m_0/m_l = \text{ratio}$ of the mass of the dissolved gas to the mass of the pure liquid in which it is dissolved.

 $f = m_q/(m_q + m_l) = r/(1+r) = \text{mass of gas per unit mass of}$ solution.

 f_q = mass of the pure gas per unit mass of the gas phase.

 $m = \rho_0 \lambda = \rho_0 f = \text{mass of gas per unit volume of solution.}$

 $x = n_g/(n_g + n_l)$ = mole fraction of the gas in the solution.

 $\delta_1 \equiv (\lambda' - \lambda)/\lambda$, where $\lambda' = \text{value of } \lambda$ that is computed from the corresponding value of r/P in Table 230, and both λ and λ' refer to P=1atm and to the indicated temperature $(t \, {}^{\circ}C)$.

P atm and p mm-Hg = partial pressure of the gas; p = 760P.

 $P_t = \text{total pressure}.$

The values of α given by Loomis have been reconverted to λ 's by multiplication by (1 + t/273.1); see text, p. 550.

Unit of P and $P_t = 1$ atm; of p = 1 mm-Hg; of r/P = 1 g of gas per atm per kg of water; of f/P = 1 g of gas per atm per g of solution; of x = 1 formula-mole per cent; of $\lambda = 1$ cm³ of gas under existing conditions per liter of solution; of $\rho_0 \lambda/P = 1$ g of gas per atm per liter of solution; of $\delta = 1$ per cent. Temperature = t °C.

Table 233.—(Continued)

Gas→		Hg Hyd	rogena	,		() ₀ ()xvg	en*	
$\stackrel{t\rightarrow}{p}$	λ (I	230	$\stackrel{t\rightarrow}{P}$	$25 \\ 1000 \ r/P$	$t \rightarrow p$	O ₂ Oxyg 23.0 λ (I	25.9 CT)4——	•
760 900 1000 1500	(17.9 ₇) 17.9 ₆ 17.9 ₆ 17.9 ₅	(17.3 ₆) 17.3 ₆ 17.3 ₅	25 50 100 200	1.57 1.56 1.55 1.52	760 900 1000 1500	(29.4 ₀) 29.3 ₉ 29.3 ₈ 29.3 ₇	(28.4 ₆) 28.4 ₆ 28.4 ₆ 28.4 ₄	
2000 2500	17.9₅ 17.9₂	17.3 ₅	400 600	1.48 1.43 ₅	2000 2500	29.3 ₄ 29.2 ₇	28.3 ₉ 28.3 ₀	
3000 3500 4000 4500	17.9 ₀ 17.8 ₆ 17.8 ₀ 17.7 ₀	17.3 _n 17.3 ₁ 17.2 ₇ 17.1 ₀	800 1000	1.40° 1:36°	3000 3500 4000 4500	29.1 ₅ 28.9 ₉ 28.8 ₁ 28.6 ₈	28.2 ₀ 28.0 ₈ 27.9 ₅ 27.8 ₁	
5000 5500 6000 6500 7000	17.5 ₈ 17.4 ₄ 17.2 ₉ 17.1 ₁ 16.9 ₃	17.0_{0} 16.9_{7} 16.8_{3} 16.6_{8} 16.5_{2}			5000 5500 6000 6500 7000	28.4 ₅ 28.2 ₄ 28.0 ₄ 27.8 ₂ 27.6 ₀	27.6 ₇ 27.5 ₁ 27.3 ₄ 27.1 ₆ 26.9 ₇	
7500 8000 Ref ⁴ δ ₁	16.7 ₀ 16.4 ₆ C 9.6	16.3 ₃ 16.1 ₂ C 11.8	Ref ^a	WGH,	7500 8000 Ref ⁴ δ ₁	27.3 ₈ 27.1 ₅ C 8.9	26.7 ₄ 26.5 ₄ C 8.1	
Gas→ t→ p	19.4 λ*(I	24.9 CT) ^d —	$P \xrightarrow{t \to P}$	N ₂ Nitroger	$\frac{1}{50}$ — $\frac{50}{1000r/P}$ (V	75 VGH ₂)d, h —	100	
760 900 1000 1500 2000	(16.2_0) 16.1_7 16.1_5 16.0_8 15.9_0	(14.9_7) 14.9_6 14.9_4 14.8_7 14.8_2	25 50 100 200 300	18.4 ₀ 16.8 ₆ 15.81 14.16 12.76	13.6 ₁ 13.3 ₂ 12.64 11.44 10.56	12.7 ₂ 12.3 ₆ 11.83 11.83 10.06	13.3 ₂ 12.9 ₆ 12.33 11.40 10.61	
2500 3000 3500 4000	15.9 ₂ 15.8 ₄ 15.7 ₆ 15.6 ₇	14.7 ₈ 14.6 ₄ 14.5 ₄ 14.4 ₆	500 800 1000 t→ P	11.10 ₆ 9.58 ₈ 8.94 0	9.30 ₄ 8.16 ₁ 7.65 ₇ 25 — 1000r/P (G	8.96 ₂ 7.91 ₁ 7.42 ₀	9.50 ₂ 8.38 ₆ 7.82 ₃	
4500 5000 5500 6000 6500 7000	15.5 ₇ 15.4 ₈ 15.3 ₇ 15.2 ₅ 15.1 ₈ 15.0 ₂	14.3 ₆ 14.2 ₇ 14.1 ₇ 14.0 ₈ 13.9 ₈ 13.8 ₉	100 125 200 300 t→	18.2 17.6 20.0 15.0	13.4 14.4 17.2 20.0	12.5 12.4 15.6 12.5	11.7 11.5 14.2 11.9	
7500 8000 Ref ^d	14.8 ₀ 14.7 ₅ C 3.1	13.7 ₆ 13.6 ₆ C 4.2	P 100 125 200 300	11.9 11.7 14.0 12.1	— 1000r/P (C 12.8 13.0 16.8 14.4	13.5 15.2 20.6 16.0		
Gas→		O	Gas→			0./		
$\stackrel{t\rightarrow}{p}$	~ λ (IC	T, C) d	$t \xrightarrow{P}$	20	ρ _θ λ/Ρ (ΙΟ	35 CT, Sa) ^d —	60	
760 900 1000 1500 2000	(27.8 _*) 27.8 ₇ 27.8 ₆ 27.8 ₄ 27.8 ₂	(27.1 ₀) 27.1 ₀ 27.1 ₇ 27.1 ₈ 27.0 ₀	25 30 35 40 45	1.29 ₀ 1.20 ₀ 1.13 ₀ 1.08 ₇ 1.05 ₁	0.66 ₀ 0.66 ₄	0.74 ₇ 0.75 ₁	0.4 ₂ 0.4 ₀	
2500 3000 3500	27.8 ₀ 27.7 ₇ 27.7 ₄	27.0 ₄ 26.9 ₈ 26.9 ₄	50 53 55	1.01 ₀ 1.00 ₀	0.66 ₈	0.75 _a	0.39 ₈ 0.39 ₁	
4000 4500	27.7° 27.6°	26.8 ₉ 26.8 ₈	60 65		0.67 . 0.67 .	0.75 ₇ 0.75 ₇	0.38 ₀ 0.38 ₀	

Table 233.—(Continued)

~	-	•	Table 233	-(Commie			
Gas→ ←	CO 17.7 λ (ICT,	19.0	$\begin{matrix} Gas \rightarrow & \frown \\ t \rightarrow & P \end{matrix}$	20	CO 35 - ρgλ/P (ICT	35	60
5000 5500 6000 6500 7000	27.6 ₁ 27.5 ₆ 27.5 ₀ 27.4 ₄ 27.3 ₇	26.7 ₈ 26.7 ₂ 26.6 ₄ 26.5 ₅ 26.4 ₈	70 80 90 100 110		78.72	0.75 _p	0.38 _b 0.39 ₅ 0.40 ₇ 0.41 ₅ 0.42 ₉
7500 8000 δ ₁	27.2 _e 27.1 ₅ -7.8	26.2 ₀ 26.1 ₅ -7.2					
Gas→	60	CO ₂	100	$Gas \rightarrow t \rightarrow t$	0	—— NH3k —	201
$\overset{t\rightarrow}{P}$		- ρ _ρ λ/P (ICT,	Sa) d	¬ • •	~	101 r/P(ICT)d	
40	0.42			300		104,	746.2
45 50	$0.4_{1} \\ 0.41_{1}$			500 700		822 714. ₇	626. ₈ 551. ₉
55	0.40			900	842	644.8	499.2
60	0.40_{0}	0.214		1000	831		_
65	0.411	0.208		1100	824	601.7	462. ₆
70	0.40_{o}	0.204	0.15_{0}	1200	816		100
80	0.40	0.199	$0.16_{7} \\ 0.17_{7}$	1300	812	571. ₉	433.,
90 100	0.41_7 0.43_1	0.199_{0} 0.199_{0}	0.177	1400 1500	80 _s 80 ₅	550.1	412.9
						330.1	712.9
110 120	0.445	$0.199_{s} \\ 0.199_{s}$	0.18 ₀ 0.19 ₀	1600 1700	$\frac{80_2}{79_5}$		
130		0.199_{3}^{3}	0.187	1800	780		
140		0.199_{1}°	0.18_{0}	Ref.d	(NP)	(Foote)	(Foote)
150		0.199 _s					
160		0.196_{8}					
Gas→		407		NH_3k			10.
$\stackrel{t\rightarrow}{p}$	20	301 - r/P (ICT)d	40	$t \rightarrow p$	O De	20 nsity; unit—:	l g/liter —
300	•	537.5	·	800		878	899
500	# 40	468.1		1000	839	868	890
700	540	421.1	312	1200	828	858	881
800 900	517 496	384.7	312	1400 1600	81 <i>7</i> 80 <i>7</i>	851 845	874 868
	478	004.7	293		798,		
1000 1100	462	358.4	293	1800 2000	190,	840 836	864 860
1200	448	0001	274.1	2200		831	856
1300	436	337.5		2400		828	852
1400	424		256.1	2600		824	848
1500	413	319.,		2800		822	844
1600	403		241.8	3000		819	841
1700 1800	392 382		229.3	3200 3400		816	837 833
1900	375			Ref.		ICT (N	
2000	365		219.3	$t \rightarrow$	25% r/P	$t \rightarrow p$	25k r/P
2100	356.1		011	p 276			
2200	347.8		211.1	3.76 6.5	1051 101 <i>7</i>	28.0 31.75	9 72 98 7
2300 2400	339. ₀ 330. ₀		203.9	9.25	1017	40.6	968
2500	326.			9.45	1007	48.5	937
2600 2600	320.4		197. _s	13.5	985	120.1	804
2700	314.4			17.19	985	173.9	769
2800	309.8		191.5	23.7	983		n
2900	304. ₅			Ref.d	Br, Hou	ı Ref.⁴	Br, Hou

Table 233.—(Continued)

Gas	,						M	JH2k						
$t \rightarrow$		20		301		40								•
Þ	_		r/	P (IC	T)d									
300	0	300.	9			185.4								
310		295.												
320		292.				180.2								
340			_			175.3								
360	Ó					170.4								
Re	f.ª	NP		Foot	e	NP								
$t\rightarrow$		25 —		30 —		35 —		45 —		55		65 —	7	′5 -
x	p	f/P	p	f/P	p	f/P	p	f/P	p	f/P	p	f/P	p	f/P
25.6	31	6.0	23	7.0	16.5	11.3	9	21						
32.7 35.9	61.5 79.5	3.9 3.3 ₁	23 45 59	5.3 4.5	31.5 43.5	7.6 6.0 ₅	9 16 21.5	15	8 11	30 24	4	60		
39.7	110	2.6_{5}	83	3.52	61	4.78	30.5	12.2 9.6	14	21	8.5	34		
51.5	271.5	1.40	207	1.84	155	2.45	83	4.59	42	9.0	19.5	19.5	13	29
54.7	335.5	1.21	257	1.58	194	2.09	106	3.82	54	7.5	25	16	17	24
62.7 66.3	597	0,83	399 469	1.17 1.05	305.5 360	$\frac{1.52}{1.38}$	172.5 205	$\frac{2.7_0}{2.41}$	92 110	5.1 4.2	45.5 55	10.2 9.0	31 38	15.1 12.9
69.5	668	0.78	525	0.98	407	1.27	233	2.23	127	4.00	65	7.9	45.5	11.4
74.6	774	0.72	610	0.91	472	1.18	274	2.04	149	3.75	77	7.3	53	10.5
77.8			650	0.90	505	1.16	295	1.98	162	3.62	84	7.0	58	10.1
84.1 88.0			736 774	0.86 0.86	573 602	$\frac{1.11}{1.10}$	334	1.90	185 196	3.42	96 102	6.6	67 72	9.5 9.2
100.0			//4	0.00	699.5	1.09	355 409.5	1.87 1.86	226	3.39 3.36	117.5	6.6 6.4 ₇	82.5	9.2
												•		

Ref. d Smits and Postma (1914), republ. by Postma (1920).

I. L. Clifford and E. Hunter (1933). The values for f/P_t have been derived from the immediately preceding values of f and P_t . t_0 and t_{100} = temperatures at which 100f = 0 and 100, respectively.

$t \rightarrow 100f$	60	80	90	100	110	120	130	140	150
0 5 10 15 20 25 30	0.197 0.439 0.717 1.084 1.559	0.467 0.90 1.48 2.19 3.05 4.14 5.55	0.692 1.30 2.00 2.91 4.12 5.55 7.12	1.000 1.82 2.75 3.88 5.31 7.18 9.48	1.414 2.41 3.58 5.03 6.91 9.20 11.98	1.96 3.22 4.70 6.55 8.83 11.72 15.46	2.67 4.27 6.15 8.42 11.33 15.04 19.8	3.57 5.55 7.83 10.70 14.37 19.20	4.70 7.14 10.02 13.58 18.6
$t \rightarrow 100f$	60	80	90	100	110 1000 f/Pt -	120	130	140	150
5 10 15 20 25 30	114.0 139.5 138.5 128.4	55.6 67.6 68.5 65.6 60.4 54.1	38.5 50.0 51.6 48.5 45.0 42.1	27.5 36.4 38.7 37.7 34.82 31.64	20.7 27.9 29.82 28.94 27.17 25.04	15.53 21.30 22.90 22.66 21.31 19.41	11.70 16.26 17.81 17.65 16.62 15.15	9.01 12.76 14.01 13.91 13.02	7.00 9.98 11.05 10.75
P_{t}		0.2	0.5	1	2 100f	4	6	8	10
17: 16: 15: 14: 13: 12: 11: 10: 9: 8: 7: 6: 5:	0 0 0 0 0 0 0 0 0 0	2.7	0.3 6.2	0 2.9 6.1 10.0 14.4 19.0	0.15 2.9 6.2 10.0 14.0 18.5 23.4 28.5	1.0 4.2 7.7 11.5 15.5 19.8 24.4 29.2 34.2 39.7	2.6 5.9 9.6 13.5 17.6 22.0 26.6 31.4 36.6 41.9 47.5	0.3 3.1 6.4 10.2 14.1 18.3 22.4 26.7 31.4 36.4 41.9 47.5 54.1	3.4 6.6 10.1 14.0 18.0 22.0 26.3 30.8 35.6 40.6 46.0 52.0 60.0

Table 233.—(Continued)

Gas→ Pi→ t	0.01	0.2	0.5	1	2 100f	4	6	8	10
40 30 20 10 0 -10 -20 -30 -40 -50 -60 -70 -72 -82.9		5.5 10.0 15.0 20.0 25.3 31.8	15.0 26.0 37.2 43.1 51.2 60.0 75.3	24.0 29.1 34.6 40.6 47.3 55.3 65.4 86.0	33.7 39.3 45.6 52.6 61.4 75.0	45. 52. 60.2 71.5 94.7	? 74.6	73.5	70.2 87.0
$t_0 \rightarrow t_{100} \rightarrow$	17.7	60.4 -61.0	81.7 46.3	100 -33.2	120.6 -18.5	144.1 -1.5	159.3 +9.7	171 18.5	180.5 25.3
t→ 100f	60	80	90	100	110 1000 fg -	120	130	140	150
5 10 15 20 25 30	564 758 841 913	478 699 812 870 908 933	485 675 771 845 891 924	456 645 753 828 879 913	418 615 737 814 862 890	393 600 727 792 832	377 582 696 768 820	347 538 672 757	313 315 ^m 664 755
t→ 100f	60	80	90	100	110 000 f ₀ /P _t .	120	130	140	150
5 10 15 20 25 30	1285 1057 776 586	531 472 371 285 219 168	373 338 265 205 160.5 129.8	250.5 234.5 194.0 156.0 122.5 96.3	173.5 171.8 146.5 117.8 93.7 74.3	122.0 127.6 111.0 89.7 71.0	88.3 94.6 82.6 67.6 54.5	62.6 68.7 62.8 52.7	43.8 31.4 ** 48.9 40.6
$P_{i} \rightarrow C$	0.2	0.5	1	2	1000 fg	4	6	8	10
170 160 150 140 130 120 110 100 90 80 70 60 50 40 30 20 10	380 655 800 905 955 980 990	45 636 875 975	318 526 700 825 902 945 970 985 992	15 290 506 675 803 892 948 975 984	3 5 6 7 8 9 9	98 30 222 63 60 36 00 48 73	195 400 571 699 784 848 900 942 970	23 207 389 5555 680 773 840 895 933 962 982	205 363 515 648 742 811 870 917 951 974 988
$t_0 \rightarrow t_{100} \rightarrow -$	60.4 -61.0	81.7 -46.3	$\frac{100}{-33.2}$	120.6 18.5		1.1 1 1.5 -	59.3 ⊦9.7	171 18.5	180.5 25.3

Table 233 .- (Continued) '

"See also Table 234.
"F. N. Speller (Sp) has given certain graphical representations of data obtained by others for the solubility of O₂ in water (ICT). Additional data have been recently reported by FTHP, dubiety = ±5 per cent.

Drucker and Moles (DM) found $\lambda = 19.6$ cm³/l for H₂ at 25 °C and total pressures of 560 to 730 mg-Hg (ICT). Additional data have been recently reported by FTHP, dubiety = ± 5 per cent, and by V. V. Ipatieff, S. J. Droujina-Artemovitsch, and V. F. Fikomiroff 106 for the ranges 20 to 140 atm and 0.5 to 45 °C.

⁴ References: See end of this section. Sets of data taken from Loomis's compilation (ICT) are each accompanied by an additional reference to its original source.

⁶ WGH₁ state that between 50 and 1000 atm their data for H₂ may be represented within their estimated error by the formula $r/\rho_0 = 24.4 + 17.2P - 0.00196P^3$ g of H₂ per kg of water, ρ_0 being the density of H₂ at 0°C and 1 atm; but that at lower pressures, the formula requires impossibly high values of r/ρ_0 .

See also Table 234 and Coste's discussion of available data. DM found for N₂ 125°C > 15.6 cm²/₁ cost the great 270 to 220°C > 15.6 cm²/₁ cost the great 270°C.

at 25 °C $\lambda = 15.6_2$ cm⁸/l over the range 270 to 830 mm-Hg (ICT). Additional data

"Nitrogen prepared chemically; not derived from air.

"Purified commercial nitrogen; 99.9 per cent N₂, the rest being argon with traces of CO₂. When P is constant, r has a minimum near 70 °C.

of CO₂. When P is constant, r has a minimum near 70 °C.

'Commercial nitrogen purified by passage over Cu at 450 °C and over soda-lime.

'Additional data have been recently reported by FTHP, dubiety = \pm 5 per cent.

'Scheffer and de Wijs (SdW) have stated that p = 0.758m (1 + 0.00270m) mm-Hg if t = 25 °C and $m \ge 2.7$ g of NH₃ per liter of solution; $m = \rho_0 \lambda$. This formula demands that r/P shall increase to a finite limit (1003) as p is indefinitely decreased. That limit is 5 per cent lower than the value found by Br for p = 3.76 mm-Hg (see elsewhere in this section of the table). The ICT gives Wijs as the source of the information here credited to SdW; the compiler has not seen Wijs's dissertation. For suggested explanations of the variation of r/P and of f/P with P, see G. Calingaert and F. E. Huggins, Jr., ¹⁰⁷ and E. Klarmann. ¹⁰⁸

'Numbers in this column are values of r/P_1 not of r/P.

"This should probably be 515, agreeing with the corresponding value under $P_1 = 10$ in the last portion of this table.

"This should probably be 51.4; see note m regarding the value from which this

This should probably be 51.4; see note m regarding the value from which this was derived.

Tables 88 with those just defined are as follows, the names used in those Tables to designate each being given after its equation:

 $\alpha = V_0/V_1P_g = (\rho_1/\rho_0)(r/P_g)$, Bunsen absorption coefficient,

 $\beta = \alpha P_a/P_t = (\rho_1/\rho_0)(r/P_t),$

 $\gamma = \alpha/\rho_l = r/\rho_0 P_g$, Kuenen absorption coefficient,

 $\delta = 100\alpha\rho_0 = 100r\rho_l/P_g$, Raoult absorption coefficient,

 $\lambda = \lambda = V_g/V_s = r\rho_s/\rho_g(1+r)$, Ostwald absorption coefficient,

 $K = p_g/x = p_g + (M_g/M_l)(\rho_l/\rho_0\alpha) = p_g(1 + M_g/M_lr)$, Henry's law constant. The unit of pressure is 1 atm for α , β , γ , δ , and 1 mm-Hg for K.

Miscellanea.

Attention must be called to a very common, but fundamentally erroneous, interpretation of the coefficients of absorption. Let $p_g = p$ mm-Hg. It is very frequently stated that $\alpha = 760V_0/V_1p$ is the volume, under

¹⁰⁰ Ipatieff, V. V., Droujina-Artemovitsch, S. J., and Fikomiroff, V. F., Jour. de Phys. (7), 3, 512D (A) (1932) ← J. chim. gén. Russe (Russian), 1, 594-597 (1931).

¹⁰⁷ Calingaert, G., and Huggins, F. E., Jr., J. Am. Chem. Soc., 45, 915-920 (1923).

¹⁰⁸ Klarmann, E., Z. anorg. allgem. chem., 132, 289-300 (1924).

Table 234.—Effect of Pressure on the Solubility of A, H₂, He, and N₂ in Water

Additional data for H₂ and N₂ are in Table 233.

 $r = m_g/m_l$ = ratio of the mass of the dissolved gas to the mass of pure water in which it is dissolved.

P = partial pressure of the gas; $P_t = \text{total pressure}$.

Unit of P and $P_t = 1$ atm; of r = 1 mg of gas per kg of water. Temp. = t °C

$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	A	rgona ——			Nitrog			
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$							$P_{t} \rightarrow t$	300 r/P t
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$							50	10.72
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	25							
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	50							
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	75							
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$								
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	100							
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	125	12.20	240.0	25.35	240	27.38	230	25.21
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	P->	25	50	Hyd	rogens	150	200	300
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	t ~				r/P			
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	0			1.919				1.839
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	10	1.751	1.742	1.741	1.736	1.721	1.706	1.672
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	20	1.617	1.608	1.607	1.604	1.587	1.572	1.545
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	30	1.533	1.524	1.523	1.518	1.503	1.488	1.467
60 1.457 1.455 1.451 1.447 1.435 1.424 1.407 1.472 1.469 1.467 1.463 1.451 1.442 1.42 80 1.511 1.507 1.503 1.498 1.489 1.477 1.45 90 1.576 1.568 1.556 1.552 1.544 1.529 1.51 1.00 1.659 1.639 1.624 1.622 1.606 1.593 1.56 1.593 1.56 1.593 1.56 1.593 1.56 1.593 1.50 1.639 1.624 1.622 1.606 1.593 1.50 1.60 1.640 1.614 1.589 1.568 1.544 1.520 1.497 1.476 1.459 1.439 1.420 1.40 1.520 1.497 1.476 1.459 1.439 1.420 1.40 1.403 1.385 1.368 1.351 1.336 1.321 1.31 1.30 1.385 1.368 1.351 1.336 1.321 1.31 1.31 1.318 1.305 1.29 1.386 1.387 1.371 1.351 1.336 1.320 1.307 1.29 70 1.404 1.387 1.368 1.351 1.336 1.320 1.307 1.29 70 1.404 1.387 1.368 1.351 1.336 1.320 1.307 1.29 70 1.404 1.387 1.368 1.351 1.336 1.320 1.307 1.29 70 1.404 1.387 1.368 1.351 1.368 1.352 1.336 1.322 1.30 80 1.436 1.417 1.397 1.381 1.367 1.352 1.33 1.37 1.461 1.448 1.424 1.411 1.393 1.37 1.00 1.537 1.515 1.497 1.478 1.458 1.437 1.41							1.443	1.422
60 1.457 1.455 1.451 1.447 1.435 1.424 1.407 1.472 1.469 1.467 1.463 1.451 1.442 1.42 1.42 80 1.511 1.507 1.503 1.498 1.489 1.477 1.45 90 1.576 1.568 1.556 1.552 1.544 1.529 1.51 00 1.659 1.639 1.624 1.622 1.606 1.593 1.56 1.593 1.56 1.593 1.56 1.593 1.50 1.624 1.622 1.606 1.593 1.56 1.593 1.50 1.624 1.622 1.606 1.593 1.50 1.624 1.622 1.606 1.593 1.50 1.50 1.640 1.614 1.589 1.568 1.544 1.520 1.49 1.640 1.614 1.589 1.568 1.544 1.520 1.49 1.403 1.385 1.368 1.351 1.336 1.321 1.31 1.30 1.445 1.424 1.407 1.389 1.373 1.356 1.34 1.403 1.385 1.368 1.351 1.336 1.321 1.31 1.50 1.386 1.368 1.351 1.334 1.318 1.305 1.29 1.386 1.387 1.371 1.351 1.336 1.320 1.307 1.29 1.404 1.387 1.368 1.351 1.336 1.320 1.307 1.29 1.404 1.387 1.368 1.352 1.336 1.322 1.30 1.307 1.29 1.404 1.387 1.368 1.352 1.336 1.322 1.30 1.307 1.29 1.404 1.387 1.368 1.352 1.336 1.322 1.30 1.307 1.29 1.483 1.461 1.448 1.424 1.411 1.393 1.37 1.351 1.390 1.483 1.461 1.448 1.424 1.411 1.393 1.37 1.351 1.360 1.537 1.515 1.497 1.478 1.458 1.437 1.41	50	1.462	1.454	1.452	1.449	1.435	1.422	1.407
70								1.406
80								1.422
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$								1.458
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$								
$P \rightarrow 0$ 400 500 600 700 800 900 100 100 0 1.800 1.768 1.742 1.717 1.687 1.653 1.61 10 1.640 1.614 1.589 1.568 1.544 1.520 1.49 20 1.520 1.497 1.476 1.459 1.439 1.420 1.40 30 1.445 1.424 1.407 1.389 1.373 1.356 1.34 40 1.403 1.385 1.368 1.351 1.336 1.321 1.31 50 1.386 1.368 1.351 1.334 1.318 1.305 1.29 60 1.387 1.371 1.351 1.336 1.320 1.307 1.29 70 1.404 1.387 1.368 1.352 1.336 1.322 1.30 80 1.436 1.417 1.397 1.381 1.367 1.352 1.33 90 1.483 1.461 1.448 1.424 1.411 1.393 1.37 90 1.483 1.461 1.448 1.424 1.411 1.393 1.37 1.515 1.497 1.478 1.458 1.437 1.41								
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$								
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		400	500	600		800	900	1000
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$		1.800	1.768	1.742		1.687	1.653	1.618
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	10	1.640	1.614	1.589	1.568	1.544	1.520	1.494
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		1.520				1.439	1.420	1.401
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$								1.342
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$								1.310
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	50	1 386	1 368	1 351	1 334	1 318	1.305	1.295
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		1 387			1 336			1.295
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$								
90 1.483 1.461 1.448 1.424 1.411 1.393 1.37 00 1.537 1.515 1.497 1.478 1.458 1.437 1.41 P \rightarrow 25 50 . 75 100 150 200 300 t 1.650 1.669 1.660 1.649 1.631 1.612 1.57 25 1.539 1.547 1.526 1.516 1.511 1.506 1.47 50 1.589 1.587 1.582 1.576 1.548 1.548 1.51					1.332	1.357		
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$								
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$								
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	00	1.55/	1.515			1.456	1.43/	1.410
0 1.650 1.669 1.660 1.649 1.631 1.612 1.57 25 1.539 1.547 1.526 1.516 1.511 1.506 1.47 50 1.589 1.587 1.582 1.576 1.548 1.548 1.51	<i>P</i> →	25	50			150	200	300
25 1.539 1.547 1.526 1.516 1.511 1.506 1.47 50 1.589 1.587 1.582 1.576 1.548 1.548 1.51	t				r/P			
50 1.589 1.587 1.582 1.576 1.548 1.548 1.51								1.573
							1.506	1.475
75 1744 1746 1720 1721 1717 1700 170			1.587	1.582				1.518
75 1.744 1.740 1.739 1.731 1.717 1.702 1.66	7 5	1.744	1.746	1.739	1.731	1.717	1.702	1.669
P→ 400 500 600 700 800 900 100		400	500	600		800	900	1000
		1.533	1.498	1.462		1.390	1.356	1.325
4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4								1.296
					1 418			1.345
								1.473

Table 234.—(Continued)

Hydrogen and nitrogen, mixture at 0 °C and 1 atm contains 76.42 vol. % of H₂. They conclude that within a few per cent each gas is dissolved as though the other were absent. $v_0 \text{ cm}^3 = \text{volume} (0 \,^{\circ}\text{C} \text{ and } 1 \text{ atm})$ of the mixed gases that is dissolved per gram of water at 25 °C; P atm = partial pressure of the hydrogen and nitrogen mixture in the gas phase: $\Delta =$ probable error of v_0 .

\boldsymbol{P}	50	100	200	400	600	800	1000
v_{o}	0.8349	1.643	3.209	6.068	. 8.809	11.327	13.724
Δ_o	0.0005	0.001	0.003	0.002	0.004	0.006	0.014
$1000 \ v_0/P$	16.70	16.43	16.04	15.17	14.68	14.16	13.72

" References:

Argon: B. Sisskind and I. Kasarnowsky.¹⁰⁰
Hydrogen: R. Wiebe and V. L. Gaddy¹¹⁰; the effect of the H₂ upon the vapor pressure of the water was considered and allowed for by the observers.

Nitrogen: A. W. Saddington and N. W. Krase.¹¹¹ They believe the numerical values reported by J. Basset and M. Dodé ¹¹² to be unreliable on account of an oiling of the surface; they have not been included in this compilation. Helium: R. Wiebe and V. L. Gaddy.^{112a}
Hydrogen and nitrogen mixture: R. Wiebe and V. L. Gaddy.¹¹⁸

For this value, t = 3 °C.

standard conditions, of the gas that "is dissolved in one volume of the solvent when the partial pressure of the gas is 760 mm," and similarly for the other coefficients. That statement is fundamentally erroneous. observations tell us only what happens when $p_q = p$ mm-Hg. Multiplying by 760 does not increase our information, but merely changes the unit in terms of which the data are expressed. Before multiplication by 760. a is the mean coefficient per mm-Hg over the range 0 to p mm-Hg; after the multiplication it is the mean coefficient per atmosphere over the range 0 to p mm-Hg, and nothing more. To infer from the observations that the latter value of α is the volume actually dissolved when $p_q = 760$ mm-Hg involves an assumption—either that p_q itself is 760 mm-Hg, or that α is independent of p throughout the range from p mm-Hg to 760 mm-Hg. Although in many cases the latter is undoubtedly true within the limits of experimental error, it is not always true, and the tacit incorporation of such assumptions into the definitions, either of the coefficients or of the symbols, is most undesirable. It introduces such confusion as may be found in International Critical Tables, vol. 3, p. 256, top of column 2, where the definition of α (p. 254) requires one to infer that at a fixed temperature the amount of hydrogen dissolved when its partial pressure is 760 mm-Hg varies all the way from 16.0 to 14.7, depending upon the conditions of observation. Of course what the data actually signify is that the mean coefficient per atmosphere is 16.0 over the range 0 to 1100 mm-Hg, and 14.7 over the range 0 to 8200. In this case it is obvious that the tacit

¹⁰⁰ Sisskind, B., and Kasarnowsky, I., Z. anorg. aligem. Chem., 200, 279-286 (1931).

 ¹¹⁰ Wiebe, R., and Gaddy, V. L., J. Am. Chem. Soc., 56, 76-79 (1934).
 111 Saddington, A. W., and Krase, N. W., Idem, 56, 353-361 (1934).
 112 Basset, J., and Dodé, M., Compt. rend., 203, 775-777 (1936).
 113a Wiebe, R., and Gaddy, V. L., J. Am. Chem. Soc., 57, 847-851 (1935).
 1148 Wiebe, R., and Gaddy, V. L., Idem, 57, 1487-1488 (1935).

Table 235.—Solubility of Atmospheric Gases in Sea-water

In the first three sections of this table are given the amounts of the indicated gases contained in a unit volume of sca-water at the indicated temperature and salinity when it is in equilibrium with air at the partial (dry) pressure of 1 atm and composed, by volume, of 79.09 per cent atmospheric nitrogen, 20.90 per cent O₂, and 0.01 per cent CO₂, the atmospheric nitrogen containing 1.185 per cent by volume of argon. They are numerically equal to the solubilities per atmosphere of partial air pressure.

In the last sections are given data for computing the "combined" CO₂ contained in sea-water in equilibrium with the air just specified. That depends upon the alkalinity (y) of the water as well as upon the partial pressure of CO₂ in the air, but is independent of the salinity. the range found in sea-water, the amount of the combined CO2 is directly proportional to y, the factor of proportionality (b) varying with the temperature and with the amount of CO₂ contained in the air.

The values in this table have been taken directly from J. J. C. Fox. 114 They agree with the tables, of different form, given by O. Krümmel 115 and based on Fox's work. Krümmel states that N₂ is essentially in equilibrium throughout the depth of the sea, but O2 is in excess near the surface, and in defect in the depths.

Fox gives the following equations, included in the compilation by D. F. Smith ¹¹⁶: For N₂, $v_0 = 18.639 - 0.4304t + 0.007453t^2 - 0.0000549t^3 C1(0.2172 - 0.00718t + 0.0000952t^2)$ cm³ per liter; for O_2 , $v_0 = 10.291 - 0.00018t$ $0.2809t + 0.006009t^2 - 0.0000632t^3 - C1(0.1161 - 0.003922t +$ $0.0000631t^2$) cm³ per liter. In each case the air is assumed to have the composition already stated, and a partial pressure = 1 atm.

J. H. Coste 117 has discussed the solution of atmospheric nitrogen in sea-water.

 $v_0 \equiv (v_t p/760) (273.1/T)$, where $v_t = \text{volume}$, at $T \circ K$ and p mm-Hg, of the gas that is contained in unit volume of sea-water when in equilibrium with air in which the partial pressure of the gas is p mm-Hg. It is the volume that v_t would occupy at 0 °C and 1 atm if the gas were ideal.

Cl = number of grams of Cl per kg of sea-water; the amount of total salts (S) per kg of sea-water is S = 1.812 Cl.

 x_0 = volume at 0 °C and 760 mm-Hg of the total amount of CO₂ contained in unit volume of sea-water; $x_0 = x_t + x_c$ where x_t is the corresponding volume of "free" CO₂, and x_0 is that of "combined" CO₂; $x_1 =$ pv_0 where p atm is the partial pressure of the CO_2 in the air, and the appropriate value of v_0 is obtained from the third section of this table:

^{*} The sign of the t^s term is incorrectly printed as + both in Fox's paper and in Smith's compilation. Fox's formula for b, not given here, also contains some typographical error.

¹¹⁴ Fox. J. J. C. Trans. Faraday Soc., 5, 68-67 (1909) ← Conseil perm. int. pour l'explor. de la mer, Publ. de Circons. No. 41, 1907; No. 44, 1909.

115 Krümmel, O., "Handbuch der Ozeanog.," Vol. 1, J. Engelhorn, Stuttgart, 1907.

116 Smith, D. F., Int. Crit. Tables, 3, 271-283 (272) (1928).

¹¹⁷ Coste, J. H., J. Phys'l Chem., 31, 81-87 (1927).

Table 235.—(Continued)

 $x_o = by$, y being the alkalinity, and the values of b being given in the last sections of this table.

y = alkalinity = number of mg OH ions per liter of sea-water completely deprived of CO_2 .

Unit of $v_0 = 1$ cm⁸ (0 °C, 1 atm) per liter of sea-water; of Cl = 1 g Cl per kg of sea-water

				Nitrog	gen			
t→ Cl	0	4	8	12	16	20	24	28
0	18.64	17.02	15.63	14.45	13.45	12.59	11.86	11.25
4	17.77	16.27	14.98	13.88	12.94	12.15	11.46	10.89
8	16.90	15.51	14.32	13.30	12.44	11.70	11.07	10.52
12	16.03	14.75	13.66	12.72	11.93	11.25	10.67	10.16
16	15.18	14.00	13.00	12.15		10.81	10.27	9.81
20	14.31	13.24	12.34	11.57	10.92	10.36	9.87	9.44
				Oxyge	en			
$t \rightarrow C$	0	4	8	12	ν ₀	20	24	28
0	10.29	9.26	8.40	7.68	-	6.57	6.14	5.75
4	9.83	8.85	8.04	7.36		6.33	5.91	5.53
4	9.36	8.45	7.69	7.04		6.07	5.67	5.31
12	8.90	8.04	7.33	6.74	6.24	5.82	5.44	5.08
16	8.43	7.64	6.97	6.43	5.96	5.56	5.20	4.86
20	7.97	7.23	6.62	6.11	5.69	5.31	4.95	4.62
		"Free"	CO ₂ , per	0.0001	atm partial	pressure		
t→ Cl c	0	4	8	12	16	20	24	28
0	0.1713	0.1473	0.1283	0.1117	$v_0 = 0.0987$	0.0877	0.0780	0.0700
2	0.1690	53	63	03	73	67	73	0.0690
4	0.1667	23	43	0.1089	59	57	66	80
6	0.1644	03	23	75	45	47	59	70
8	0.1621	0 1383	03	61	31	37	52	60
10	0.1598	63	0.1183	47	17	27	45	50
12	0.1575	43	63	33	03	17	38	40
14	0.1552	23	43	19	0.0889	07	31	30
16	0.1529	03	23	05	75	0.0797	24	20
18	0.1506	0.1283	03	0.0991	61	87	17	10
20	0.1483	0.1263	0.1083	0.0977	0.0847	0.0777	0.0710	0.0600

"Combined" CO2

For combined CO., $x_c = by$ cm³ (0 °C, 760 mm-Hg) per liter of sca-water, the unit of y being that stated at head of table; p is partial pressure of CO₂ in the overlying air, unit = 0.0001 atm.

$t \rightarrow$	0	2	4	6	8	10	12	14
ħ								
1.5	1.1673	1.1562	1.1451	1.1340	1.1229	1.1118	1.1007	1.0896
2.0	. 2021	920	818	717	615	513	412	1.1310
2.2	2140	1.2042	1.1943	845	746	647	549	450
2.4	2249	153	1.2051	1.1961	865	769	673	577
2.6	348	259	160	1.2066	1.1973	879	785	691
2.8	437	346	254	162	1.2070	1.1978	887	795
3.0	518	428	338	248	158	1.2068	1.1978	888
3.2	592	504	415	327	238	150	1.2062	1.1973
3.4	658	571	484	397	310	223	136	1.2049
3.6	718	633	547	461	376	290	205	119
3.8	773	689	603	519	434	350	265	180
4.0	822	739	655	572	488	405	322	238

			"(Combined"	CO ₂			
$t \rightarrow p$	0	2	4	6 	8	10	12	14
4.5	1.2928	848	767	687	606	525	445	364
5.0	1.3019	1.2942	864	787	709	632	555	477
5.5	105	1.3032	1.2958	885	812	739	666	592
6.0	197	130	1.3062	1.2995	1.2928	1.2861	794	726
6.5	1.3306	1.3246	1.3187	1.3127	1.3067	1.3008	1.2948	1.2889
t→ p ~	16	18	20	22 h	24	26	28	_
1.5	1.0785	1.0674	1.0563	1.0452	1.0341	1.0230	1.0119	•
2.0	1.1209	1.1107	1.1005	1.0904	802	701	599	
2.2	352	253	154	1.1056	1.0957	1.0859	760	
2.4	481	385	289	193	1.1097	1.1001	1.0905	
2.6	597	504	410	316	222	128	1.1035	
2.8	703	611	519	428	336	244	152	
3.0	798	708	618	528	438	348	258	
3.2	885	796	708	620	531	443	354	
3.4	1.1962	875	788	701	614	527	440	
3.6	1.2033	1.1948	862	777	691	605	520	
3.8	096	1.2011	927	842	757	673	588	
4.0	155	071	1.1988	1.1905	821	738	654	
4.5	284	203	1.2122	1.2042	1.1961	1.1881	800	
5.0	400	322	245	168	1.2090	1.2013	1,1935	
5.5	519	446	373	300	226	153	1,2080	
6.0 6.5	659 1.2829	592 1.2769	525 1.2710	458 1.2650	390 1.2591	$\frac{323}{1.2531}$	256 1,2471	

Table 235.—(Continued)

assumptions written into the definition do not apply to the case in hand. Such confusion is not peculiar to the *International Critical Tables*, but is to be found throughout the literature treating of the absorption of gases, and in most compilations of such data.

From the relations connecting α and λ with r it is obvious that $\alpha = (\rho_l \rho_g / \rho_b \rho_0) \{ (1+r)\lambda / P_g \}$. If the amount of gas dissolved is small, then the density (ρ_s) of the solution is essentially the same as that (ρ_l) of the pure liquid, and (1+r) is essentially unity. Under such conditions $\alpha = \rho_g \lambda / \rho_0 P_g$, essentially. Furthermore, if the gas were ideal, then $\rho_g / \rho_0 P_g$ would equal 273.1/(273.1+t) and we could write $\alpha = (273.1)\lambda / (273.1+t)$, which is the relation used by A. G. Loomis ⁸⁹ in reducing to α the data which were initially expressed in terms of λ . Actually the gases are not ideal, and in certain cases their departure from that condition must be considered.

For that reason, data so converted by him have, for this compilation, been reconverted to λ by multiplication by (1+t/273.1), his interpolations being accepted without question. The data have been extrapolated, when necessary, to $P_g=1$ atm, so as to obtain comparisons with the corresponding values computed from those of r/P_g obtained by others. In some cases marked differences exist.

The supersaturation of liquids with gases has been studied by

J. Metschl,⁹⁰ certain errors that frequently occur in the measuring of the solubility of gases have been discussed by W. Manchot,⁹¹ and the effect of colloids and of fine suspensions upon the solubility of gases has been investigated by A. Findlay and associates.⁹²

For rates of solution and of aeration of quiescent columns of water, see Section 86.

References.

In this list are given those references which have been indicated at some place in the section solely by the name of the author or by a symbol. The names and symbols are listed in alphabetical order.

In certain tables there are two kinds of references, designated as Ref₁ and Ref₂, respectively. The first includes those from which the values were obtained; the second, those that should also be considered by one especially interested. In ICT, these Ref₂ references followed the instruction "cf. also."

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   \mathbf{D}\mathbf{M}
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86. RATE OF SOLUTION OF GASES IN WATER

When a quiescent column of water, whether fresh or salt, is exposed to a gas, there is, in general, a vertical streaming which mixes the upper layers with the lower ones. This occurs even when the gas is initially saturated with water-vapor and the entire system is maintained at a uniform and contant temperature, and is much more pronounced when there is evaporation, with its attendant cooling and enhanced concentration of the surface layer. See W. E. Adeney and H. G. Becker, ¹¹⁸ W. E. Adeney, A. G. G. Leonard, and A. Richardson, ¹¹⁹ H. G. Becker and E. F. Pearson, ¹²⁰ and W. E. Adeney.

The streaming is more uniform and rapid in a salt solution than in pure water, is more rapid above 10 °C than below, and varies with the concentration of the salt. For solutions of NaCl the streaming is most rapid when the concentration is about 1 per cent. It extends to a depth of at least 10 feet.¹¹⁹

Data on the progressive aeration of quiescent and initially air-free water are given in Table 236.

When a liquid is kept thoroughly mixed while gas is entering it through a fixed surface, the net rate at which the gas enters the liquid is $dm/d\tau = (\alpha P - \beta c)A$, where dm is the amount entering through an area A in the time $d\tau$, P = partial pressure of the gas, c is the concentration of the gas in the liquid, and α and β are two coefficients; α may be called the entrance, and β the exit, coefficient. If the volume of the liquid is V and the total amount of gas in it is m, then c = m/V, and the equation may be written either $dm/d\tau = (\alpha P - \beta m/V)A$ or $dc/d\tau = (\alpha P - \beta c)(A/V)$. If the subscripts 0 and ∞ indicate that the values correspond to $\tau = 0$ and to $\tau = \infty$, respectively, i.e., the initial and the saturation values, then $c - c_0 = (c_\infty - c_0)(1 - e^{-\beta A\tau/V})$; calling $c - c_0 \equiv \Delta c$, this becomes $\Delta c/c_\infty = [(c_\infty - c_0)/c_\infty](1 - e^{-\beta A\tau/V})$. The equation still holds good if all the c's are replaced by m's. Also, the initial rate of solution when $c_0 = 0$ is $(dc/d\tau)_0 = \alpha PA/V$, $(dm/d\tau)_0 = \alpha PA$; at saturation, $(dc/d\tau) = 0$, $c = c_\infty$,

¹¹⁸ Adeney, W. E., and Becker, H. G., Sci. Proc. Roy. Dublin Soc. (N. S.), 16, 143-152 (1920) = Phil. Mag. (6), 42, 87-96 (1921).

Adeney, W. E., Leonard, A. G. G., and Richardson, A., Sci. Proc. Roy. Dublin Soc. (N. S.),
 17, 19-28 (1922) = Phil. Mag. (6), 45, 835-845 (1923).
 Becker, H. G., and Pearson, E. F., Sci. Proc. Roy. Dublin Soc. (N. S.), 17, 197-200 (1923).

¹²¹ Adeney, W. E., Idem, 18, 211-217 (1926) = Phil. Mag. (7), 2, 1140-1148 (1926).

Table 236.—Aeration of Quiescent Water 126

The water was quiescent and initially air-free. v_0 = volume (at 0 °C and 760 mm-Hg) of air per liter of water at a point 12 cm below the free surface; v_s = value of v_0 at saturation, when $\tau = \infty$; $\tau =$ duration of the exposure: temperature = 13.5 °C.

The formula $(v_8 - v)/v_8 = e^{-k\tau}$ will not fit the observations unless k is a function of τ , decreasing at an ever decreasing rate as τ increases.

With reference to the solution by water of CO₂ from the atmosphere, J. Johnston 127 has stated that "even in an unstirred liquid contained in an open beaker the process is substantially complete in about ten minutes."

		Unit of T	= 1 hr; o	$f v_0 = 1 \text{ m}$	l/liter; of v	$v_s = 1\%$		
τ	v_0	v_0/v_s	τ	v_0	v_0/v_s	τ	v_0	v_0/v_x
2	5.0	21.4	20	11.6	49.6	70	16.0	68.4
4	6.3	26.9	25	12.4	53.0	80	16.4	70.1
6	7.5	32.0	30	13.2	56.4	90	16.9	72.2
8	8.2	35.0	40	13.9	59.4	100	17.4	74.3
10	8.9	38.0	50	14.7	62.8	00	23.4	100.0
15	10.3	44.0	60	15.4	65.8			

Table 237.—Entrance Coefficient of Gases into Water

The entrance coefficient is the α occurring in the expression $dm/d\tau =$ $(\alpha P - \beta c)A$; whence $\alpha = \beta m_{\infty}/PV$; m_{∞}/PV is a mean coefficient of absorption between 0 and P, write $s \equiv m_{\infty}/PV$. The following values of βs have been computed from the values of β in Table 238 and those of s $(\equiv \rho_l f/P)$ in Tables 230 and 231, ρ_l being the density of the solution; for our present purposes, ρ_l may be taken as 1 g/cm³, and f/P as r/P. It will be noticed that α and βs are nearly independent of the temperature.

	Unit	of a and	$\beta s = 10^{-3} \text{m}$	g/min.cm ² a	tm. Temp.	$= t \circ C$	
Gas→	O_2	N_2	CO ₂	Air	Gas→	Bs CC	Ω_{2}^{a}
o ´	24.3	9.9	256	12.5	0.1	256	245
5	23.7	10.1	266	12.7	9.9	266	249
10	24.2	10.3	268	12.8	22	259	33 %
20	24.6	10.8	262	12.9	34.6	245	217
30	24.6	11.2	253	13.2	43.1	234	219
40	24.6	11.7	242	13.6			

^a From C. Bohr ¹²² excepting at 22 °C.
^b From H. G. Becker ¹²⁴ who gives, among others, the following approximate values for the initial rates of solution at 22 °C. It is believed that the volumes were measured at 22 °C and under the pressures (P) at which the absorption occurred, and it is on this assumption that the values of P here tabulated were computed from his two sets of rates. The value of α has been taken as (rate/P). It will be noticed that his value for CO₂ is entirely inconsistent with those of Bohr.

Gas	H_2	CO ₂	Cl2	H_2S	SO ₂	HCl	
Rate	0.0043	0.0168	0.0984	0.0950	2.00	54.3	cc/min·cm ²
Rate	0.000023	0.000198	0.00187	0.000866	0.0343	5.3	g/hr·cm²
P	0.991	0.0994	0.0986	0.0984	0.0967	0.992	atm
α	0.00038	0.033	0.316	0.146	5.90	89.	mg/min·cm³-atm

¹²² Bohr, C., Ann. d. Physik (Wied.), 68, 500-525 (1899).

¹²⁰ Adeney, W. E., and Becker, H. G., Sci. Proc. Roy. Dublin Soc. (N. S.), 15, 385-404, 609-628 (1918-1919) = Phil. Mag. (6), 38, 317-337 (1919); 39, 385-404 (1920).

Table 238.—Exit Coefficient of Gases from Water

The exit coefficient is the β occurring in the expression $c - c_0 =$ $(c_{\infty} - c_0)(1 - e^{-\beta A \tau/V})$ (see text). $\beta = \beta'(T - T') = \beta'(t - t')$; the tabulated values of β have been computed from the tabulated values of β' and T', which were published in the articles cited; it has been assumed that T = 273 + t. The value of β for air from sea-water is about 5 per cent less than that for air from distilled water. 123, first paper

	Unit of	$\beta = 1$ cm/min.	Temp. = $t {}^{\circ}C$	
Gas→	O ₂	N_2	CO_2	Air
10 'β'	96	103	38.1	99
T'	237	240	253	239
t'	-36	-33	-20	-34
*				
0	0.346	0.340	0.076	0.336
0 5	0.384	0.391	0.095	0.386
10	0.442	0.443	0.114	0.436
20	0.538	0.546	0.152	0.534
30	0.634	0,649	0.190	0.634
40	0.730	0.752	0.229	0.733
Ref.a	AB	ΛB	\mathbf{B}	AB

A. Guyer and B. Tobler 128 have reported the following values. Temperature of the bulk of the liquid was 20 °C.

Gas	β
Acetylene	4.32
CO ₂	3.84
H_2S	3.76
SO ₂	0.93
NHs	0.09

Table 239.—Absorption of Oxygen by a Thin Film of Water 121

It is assumed that the water is initially free of O_2 . If c = concentration at the time τ , $c_{\infty} = c$ when $\tau = \infty$, A = area of the water surface, V =volume of the film, and $\beta = 0.0096(t + 36)$ cm/min, then $c/c_{\infty} =$ $1 - e^{-\beta A\tau/V}$. The values of τ tabulated below correspond to V/A = 0.05cm: hence if the absorption occurs on one side only and there is no loss from the other side (a thin film of water on a non-porous solid upon which the gas is not adsorbed), the film is 0.05 cm thick, but if absorption occurs on both sides, the thickness is 0.10 cm. The formula assumes that there is no evaporation, that the film does not mix with any other liquid, that β and c_{∞} have the same values for the film as for an extended volume of thoroughly mixed water, and that the concentration is the same throughout the entire volume of the film. When appropriate values are given to c_{∞} , the tabulated values apply to atmospheric oxygen as well as to pure oxygen, and to sea-water as well as to pure water.

AB Adeney, W. E., and Becker, H. G. 123, second paper B Bohr, C. 122

Unit of	$\tau = 1 \text{ sec};$	of $c/c_{\infty} = 1$	%; temp. = 1	°C; V/A =	0.05 cm	
0	5	10	15	20	25	30
1.0	0.8	0.8	0.7	0.7	0.6	0.5
2.1	1.8	1.6	1.4	1.3	1.3	1.2
3.2	2.7	2.3	2.0	1.8	1.7	1.6
4.5	4.0	3 .6	3.3	2.9	2.7	2.4
6.0	5.2	4.6	4.2	3.9	3.6	3.3
8.0	7.2	6.4	5.7	5.0	4.6	4.3
10.5	9.3	8.2	7.3	6.7	6.1	5.8
14.0	12.5	11.0	10.0	9.0	8.3	7.7

14.2

28.4

13.0

26.0

11.9

23.8

10.8

21.6

Table 239.—(Continued)

15.6

31.2

and $\alpha = \beta c_{\infty}/P = \beta m_{\infty}/PV$; c_{∞}/P and m_{∞}/PV are mean coefficients of absorption between 0 and P. See C. Bohr, ¹²² W. E. Adeney and H. G. Becker, ¹²³ H. G. Becker, ¹²⁴ and W. E. Adeney. ¹²¹

For values of α and β , see Tables 237 and 238.

90

99

19.8

39.6

17.5

35.0

According to L. A. Klufschareff,¹²⁵ a superficial layer of machine oil does not affect the rate of solution of CO₂ into water, but colloid particles, whether acid or neutral, increase both the rate and the amount of the solution of CO₂.

87. DIFFUSION OF GASES IN WATER

The diffusivity, or coefficient of diffusion, of a solute in a liquid is the quantity D in the equation $(dq/d\tau)_x = -D(dc/dx)_\tau dy dz$, where dq is the amount of the solute which in the time $d\tau$ passes in the positive direction of x through the area dydz at x, the concentration at x increasing in the direction of x at the rate $(dc/dx)_\tau$; unless the contrary is stated, both q and c are specified in terms of the same unit amount of solute, and the unit of volume involved in c is the cube of the unit of length appearing in dx, dy, and dz. Whence, $(dc/d\tau)_x = D(d^2c/dx^2)_\tau$ if D is independent of x; in general, D depends upon c, and hence, upon x. In that case $(dc/d\tau)_x = \{d[Ddc/dx]/dx\}_\tau = D(d^2c/dx^2)_\tau + (dc/dx)_\tau^2(dD/dc)$. The temperature is supposed to be uniform and constant.

[&]quot;In the paper here cited this unit is incorrectly given as "per sec"; cf. with tables and graphs in AB reference of Table 238.

¹²⁴ Becker, H. G., Ind. Eng. Chem., 16, 1220-1224 (1924).

¹²⁵ Klufschareff, L. A., Jour. de Phys. (7), 3, 513D (1932) ← Jour. chim. appl. Russe, 4, 425-428 (1931).

¹²⁰ Chappuis, P., Trav. et mém. bur. int. poids et mes., 14, (B) 1-163, and D.1-D.63 (1910). ¹²⁷ Johnston, J., J. Am. Chem. Soc., 38, 947-975 (1916).

¹²⁸ Guyer, A., and Tobler, B., Helv. Chim. Acta, 17, 550-555 (1934). See also, Idem, 17, 257-271 (1934).

Table 240.—Diffusivities of Selected Gases in Water

Adapted from the compilation of H. R. Bruins. 129

The data for NH_3 represent a mean value of the diffusivity (D) over an ill-defined range of concentration, the diffusion having been from a solution of initial concentration c into water initially NH₃-free. The others refer to exceedingly dilute solutions.

		Unit of	$D = 10^{-5} \text{ cm}^2/$	sec; of c	= 1 gfw	per lit	er. Tem	p. = t °(C	
Gas	t	D	Ref. 1ª	Ref.2ª	Gas	ŧ	c	D	Ref.1ª	Ref.20
H_2	10	4.3	Hu8	Ba	NH_3	5	0.7	1.2.	Sch	$\mathbf{E}\mathbf{x}$
	16	4.7	Hu 7	Ex		5	3.5	1.24	Sch	Ha
	21	5.2	Hu 7	Ha		8	1.0	1.36	Ar	Mu
N_2	22	2.0_{2}	Hu 7	$\mathbf{E}\mathbf{x}$		8	Sat.b	1.0_{8}	Hu8	Vo
Rn	18	1.140	\mathbf{Ro}	Wa		10	Sat.b	1.14	Hu8	
CO_2	10	1.4	Hu 7, 8, St	$\mathbf{E}\mathbf{x}$		12	1.0	1.64	Ar	
	15	1.6	Hu 7, 8, St	Ha		15	1.0	1.77	Ab	
	18	1.71 4	Ca			15	Sat.b	1.26	Hu8	
	20	1.77	Hu 7. 8. St	Wr						

H₂ W. W. Ipatieff, Jr., W. P. Teodorovitsch, and S. I. Druschina-Antemovitsch ¹³⁰⁰ have reported that, within the limits 35 to 100 atm, D for H2 is independent of the pressure and has the following values:

t	15	25	35	45	° C
$10^{i}D$	2.49	3.37	4.22	5.69	cm ² /sec.

^a References: Under Ref., are the references from which the associated data were obtained; under Ref. are supplementary references pertinent to the gase but not necessarily to the particular values of t and c given in this table.

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to the particular values of t and c given in this table.

Abegg, R., Z. physik. Chem., 11, 248-264 (1893).

Arrhenius, S., Idem, 10, 51-95 (1892).

Barus, C., Carnegie Inst. of Washington, Publ. No. 186, 1-88 (1913).

Carlson, T., J. Am. Chem. Soc., 33, 1027-1032 (1911); Medd. f. K. I'ctensk. Nobelinst., 2, No. 6 (1913).

Exner, F., Ann. d. Physik (Pogg.), 155, 321-336, 443-464 (1875).

Hagenbach, A., Idem (Wied), 65, 673-706 (1898).

Hufner, G., Idem, 60, 134-168 (1897).

Hufner, G., Z. physik. Chem., 27, 227-249 (1898).

Muller, J., Ann. d. Physik. (Wied.), 43, 554-567 (1891).

Rona, E., Z. physik. Chem., 92, 213-218 (1917).

Scheffer, J. D. R., Idem, 2, 390-404 (1888).

Stefan, J., Sitab. Akad. d. Wiss. Wien (Abt. II), 77, 371-409 (1878).

Voigtlander, F., Z. physik. Chem., 3, 316-335 (1889).

Vallstabe, F., Physik. Z., 4, 721-722 (1903).

v. Wroblewski, S., Ann. d. Physik. (Wied.), 2, 481-513 (1877); 4, 268-277 (1878); 7, 11-23 (1879); 8, 29-52 (1879).
Ab
Ar
Ba
Ca
```

Ha Hu7

Hu8

Mu

Ro Sch St Vo Wa Wr

PRESSURE-VOLUME-TEMPERATURE ASSOCIATIONS FOR SATURATED WATER AND STEAM

In this section are assembled those pressure-volume-temperature associations that are characteristic of water in equilibrium with its own pure vapor, and of water-vapor in equilibrium with pure water. See also Table 260.

When such equilibrium exists, each phase is said to be saturated with

^b Diffusion from water saturated with NH₂ at a pressure of 1 atm.

 $^{^{}d}$ 1.7, \pm 0.03. $^{\circ}1.1_{4} \pm 0.07$.

¹²⁹ Bruins, H. R., Int. Crit. Tables, 5, 63-76 (1929).

¹³⁰ Ipatieff, W. W., Ir., Teodorovitsch, W. P., and Druschma-Antemovitsch, S. I., Z. anorg. allgem. Chem., 216, 66-74 (1933).

reference to the other, and its state is more briefly described by the single word "saturated," as in "saturated water," "saturated steam." *

Critical Data.

The critical data of a substance are those that are characteristic of it at its critical point. And the critical point in the usual p-v diagram is defined by the associated temperature and pressure that marks the vertex of the curve that bounds the area in which both phases (liquid and vapor) may coexist and each may be distinctly segregated. There $(\partial p/\partial v)_t = 0$ and $(\partial v/\partial t)_p = \infty$. At that point the distinction between the two phases vanishes. Consequently, the meniscus separating the phases vanishes, and so does the latent heat of transition. Each of these phenomena has been used as a criterion for determining when the critical point has been attained; the first, the more frequently.

But things are not quite so simple as one might infer. For example, E. Schröer ¹³² has stated that the densities of the two phases do not become equal as the system passes through the critical point, although the meniscus vanishes; though mixed by stirring, segregation occurs when the stirring is stopped. Only after the temperature has been raised to about 0.5 °C above the critical point does the system become homogeneous. Similar observations have been reported in some detail by O. Maass and A. L. Geddes.¹³³ who state that a difference in densities of the upper and the lower layers persisted after 6 hours of continuous stirring. This difference, at temperatures above the critical, decreases with rise in temperature, ultimately vanishing. Also, the difference is destroyed if the system is expanded while slightly above the critical temperature; and also if the upper portion of the system is cooled below the temperature of the lower one. Once the difference of density is destroyed, the system remains homogeneous however the temperature and pressure may be varied, provided that the temperature does not fall below the critical one. Such differences in

^{*} Over 50 years ago, W. Ramsay and S. Young ist proposed that the "curve representing the relations between volumes and temperatures at the vapor-pressures corresponding to the temperatures" be called the "orthobaric" curve. This has led to an occasional use of the adjective orthobaric to modify the name of a property for the occasional use of the adjective orthobaric to modify the name of a property for the purpose of indicating that the values considered are solely those characteristic of a given liquid when subjected to the equilibrium pressure of its own pure vapor; as in "orthobaric density." But the word does not appear in any of the general English dictionaries; and its significance, as defined by Ramsay and Young, is unknown to many who are interested in the data for saturated liquids, and is not at all self-evident. There seems to be little reason for using it, especially as the same purpose can be served by qualifying the name of the liquid or its class by the adjective "saturated," in accordance with a custom long established for vapors. However, should an adjective applicable to the property, instead of the substance, be needed, then either autopiestic or, less appropriately, autobaric would be preferable to orthobaric, as the first component—"auto"—directly suggests that the pressure in mind arises from, and is characteristic of, the substance itself, while "ortho" does not.

Ramsay W., and Young, S., Phil. Trans., 177, 123-156 (135-136) (1886).

¹⁸¹ Ramsay, W., and Young, S., Phil. Trans., 177, 123-156 (135-136) (1886).

¹⁸² Schröer, E., Z. physik. Chem., 129, 79-110 (1927).

¹⁸⁸ Maass, O., and Geddes, A. L., Phil. Trans. (A), 236, 303-332 (1937).

density had previously been observed by G. Teichner ¹³⁴; see also J. Traube, ¹⁸⁵ who developed a theory based on the idea that the liquid molecule differed from the gas one in size but not in mass, and that at the critical point the two types of molecules were miscible in all proportions, becoming identical only at a higher temperature. Maass and Geddes refer to other papers bearing on the subject.

Furthermore, C. A. Winkler and O. Maass ¹³⁶ have reported observations indicating that the surface energy does not become zero when the meniscus vanishes.

And H. L. Callendar ¹³⁷ has reported that differences in density can be detected as far as 6 °C above the critical temperature (374 °C), that the latent heat of vaporization does not vanish until that higher temperature is reached, and that small amounts of air in the water cause serious difficulties. These conclusions were not entirely borne out by the work of A. Egerton and G. S. Callendar ¹³⁸; and the careful work of W. Koch, ¹³⁹ specially

Table 241.—Critical Constants of Water

The values preferred at present by the National Bureau of Standards are these:

 $t_c = 374.15 \, ^{\circ}\text{C}, \ p_c = 225.65 \, \text{kg*/cm}^2 = 218.39 \, \text{atm}, \ v_c* = 3.1 \, \text{cm}^3/\text{g.}^{142}$

The several sets of values reported since 1900, excepting H. L. Callendar's (see text), are given below. The assigning of two values to a given reference indicates that the authors state no more than that the true value lies between those limits.

Unit→ Year	1 °C	1 atm	1 kg*/cm²	1 cm ³ /g	Ref^a
1904	374			•••	TT
1910	374.07; 374.62	218	225		HB
1927	374.20 ± 0.20			• • •	S
1928	374.0	217.72	224.95	2.5	ICT
1931	374.11	218.53	225.79	3.086	KS
1932	374: 374.5				EC
1932	374	219	226	2.8; 3.0	NB
1934	374.11 +	218.167	225.416		SKG
1937	374.23	218.26	225.51	3.066	E
1937	374.15	218.39	225.65	3.1	OSG

 $1 \text{ atm} = 1.03323 \text{ kg*/cm}^2 = 1.01325 \text{ bars}$

For the viscosity at the critical point, see p. 66.

E. H. Riesenfeld and T. L. Chang 143 have given for ordinary water

¹⁸⁴ Teichner, G., Ann. d. Physik (4), 13, 595-610, 611-619 (1904) = Diss., Wien, 1903.

¹⁸⁵ Traube, J., Z. anorg. Chem., 37, 225-242 (1903).

¹⁸⁶ Winkler, C. A., and Maass, O., Can. J. Res., 9, 65-79 (1933).

¹⁸⁷ Callendar, H. L., Proc. Roy. Soc. (London) (A), 120, 460-472 (1928); Engineering (London), 126, 594-595, 625-627, 671-673 (1928).

¹³⁸ Egerton, A., and Callendar, G. S., Phil. Trans. (A), 231, 147-205 (1932).

¹³⁰ Koch, W., Forsch. Gebiete Ingenieurw., 3, 189-192 (1932).

¹⁴⁰ Jakob, M., Physik. Z., 36, 413-414 (1935).

¹⁴¹ Harand, J., Monatsh. Chem., 65, 153-184 (1934).

Table 241.—(Continued)

the following values in the neighborhood of t_0 for the density (ρ) of saturated vapor and of saturated liquid. The mean of the two densities corresponding to the same t is almost linear in t, and by extrapolation leads to $\rho_c = 0.329 \text{ g/cm}^3$, or $v_c^* = 3.04 \text{ cm}^3/\text{g}$. As tabulated, the values proceed in regular order from vapor saturated at 364.0 °C to liquid saturated at 365.4 °C. Unit of $\rho = 1 \text{ g/cm}^3$; of $v^* = 1 \text{ cm}^3/g$; temp. = t °C.

ŧ	ρ	v*	1	P	v*
364.0	0.157	6.37	373.9	0.392	2.55
373.1	0.244	4.10	373.7	0.398	2.51
374.0	0.271	3.69	373.4	0.402	2.49
374.0	0.288	3.47	372.9	0.409	2.44
374.0	0.317	3.15	373.4	0.410	2.44
374.3	0.322	3.10	371.8	0.440	2.27
374.2	0.329	3.04	367.2	0.478	2.09
374.2	0.343	2.92	366.5	0.481	2.08
374.2	0.344	2.91	365.4	0.490	2.04
374.2	0.354	2.82	365.4	0.491	2.04

a References.

- EC
- Egerton, A., and Callendar, G. S. 128 Eck, H., Tatigkeit d. Phys. Techn. Reichs., p. 32, 1936, = Physik. Z., 38, 256 (1937). Holborn, L., and Baumann, A., Ann. d. Phys. (4), 31, 945-970 (1910). Int. Crit. Tables, 3, 248 (1928). Compilation by A. F. O. Germann and S. F. Pick-HB
- KS NB
- ering.
 Keyes, F. G., and Smith, L. B., Mech. Eng., 53, 132-135 (1931).
 v. Nieuwenburg, C. J., and Blumendal, H. B., Rec. Trav. chim. Pays-Bas, 51, 707-714 (1932).
 Osborne, N. S., Stimson, H. F., and Ginnings, D. C., J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937).
 Schröer, E., Z., physik. Chem., 129, 79-110 (1927).
 Smith, L. B., Keyes, F. G., and Gerry, H. T., Proc. Am. Acad. Arts Sci., 69, 137-168 (1934).
 Traube, J., and Teichner, G., Ann. d. Phys. (4), 13, 620-621 (1904). OSG
- SKG

undertaken for their investigation, has given no confirmation of them, not even of the serious effects caused by a small admixture of air. One would be inclined to think that Callendar's conclusions are erroneous were it not for the other observations that have been mentioned, but they, and especially those of Maass and Geddes, suggest that the failure in confirmation may have resulted from some difference in the procedures followed.

See also M. Takob 140 and J. Harand 141; the latter observed the region of the meniscus with a "schlieren" microscope and found the fluid to be nonhomogeneous immediately after the meniscus had vanished.

Saturated Vapor.

(a) Vapor-pressure.—Definition.—The pressure of a vapor that is in equilibrium with its liquid depends not only upon the temperature, but also upon the shape of the liquid surface and upon the presence or absence of a foreign gas, the effect of such gas varying with its partial pressure. That particular pressure of the vapor, unmixed with a foreign gas, that is in equilibrium with a flat surface of its own pure liquid is properly called the

¹⁴² Osborne, N. S., Stimson, H. F., and Ginnings, D. C., J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937).

¹⁴⁸ Riesenfeld, E. H., and Chang, T. L., Z. phys. Chem. (B), 30, 61-68 (1935).

vapor-pressure of the liquid, and is presumably the quantity that is given in tables of vapor-pressures. The equilibrial pressure under other conditions, though quite frequently called by the same name is preferably designated in a different manner.

Past History, Effect of.—Menzies and his associates have concluded from their own observations that, when the temperature of water is changed, any accompanying change in the internal state of the water is completely attained practically instantly.¹⁴⁴

Furthermore, as certain observations by T. C. Barnes and his associates ¹⁴⁵ have indicated that water from freshly melted ice is more conducive to the growth of certain biological organisms than is either ordinary water or water from freshly condensed steam, and this has been interpreted as arising from the presence of an excess of trihydrol ¹⁴⁶ in the freshly melted ice, A. W. C. Menzies ¹⁴⁷ has compared the vapor-pressure of water from freshly melted ice with that from freshly condensed steam, finding no difference. He cannot accept the trihydrol explanation (see also A. W. C. Menzies ¹⁴⁸).

Catalysts, Effect of.—Although II. B. Baker ¹⁴⁰ has reported a progressive change in the vapor-pressure of water in contact with such catalysts as charcoal, thoria, and platinum black, Menzies and his associates ¹⁵⁰ seem to have shown that Baker's conclusions cannot be accepted.

Gas, Effect of.—The effect of an inert gas upon the equilibrial pressure of the vapor of a liquid in contact with it is twofold: (1) the direct

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Table 242.—Vapor-pressure of Water (atm): -5 to +374 °C <sup>156</sup> (See also International Skeleton Steam Tables, 1934, (Table 260) and Tables 243, 244, and 245.)
```

These values, computed by means of formula (1) are believed to be the most accurate available for $t \equiv 100$ °C, for which range they were chosen for the International Skeleton Steam Tables of 1934 (see Table 260); for lower temperatures the slightly different values given in the latter tables are to be preferred. Their differences from those here given are shown in Table 246. All values refer to a flat surface of pure water in contact with its own pure vapor. For the effect of inert gases, see text (p. 560) and Tables 243 and 245.

In the Osborne-Meyers paper the pressures are expressed in kg*/cm² as well as in atm, and are given for each 1 °C; also in lb*/in² for each 1 °F.

¹⁴⁴ West, W. A., and Menzies, A. C. W., J. Phys'l Chem., 33, 1893-1896 (1929).

145 Barnes, T. C., Proc. Nat. Acad. Sci., 18, 136-137 (1932); Lloyd, F. E., and Barnes, T. C., Idem, 18, 422-427 (1932); Barnes, T. C., and Jahn, T. L., Idem, 19, 638-640 (1933); Barnes, T. C., and Larson, E. J., J. Am. Chem. Soc., 55, 50-59 (1933); Barnes, T. C., and Jahn, T. L., Quart. Rev. Biol., 9, 292-341 (1934); Barnes, T. C., Science, 79, 455-457 (1934); 81, 200-201 (1935).

146 Barnes, H. T., and T. C., Nature, 129, 691 (1932).

147 Menzies, A. W. C., Proc. Nat. Acad. Sci., 18, 567-568 (1932).

148 Menzies, A. W. C., Science (N. S.), 80, 72-73 (1934).

149 Baker, H. B., J. Chem. Soc. (London), 1927, 949-958 (1927).

¹⁵⁰ West, W. A., and Menzies, A. W. C., *J. Phys'l Chem.*, 33, 1893-1896 (1929); Wright, S. L., and Menzies, A. W. C., *J. Am. Chem. Soc.*, 52, 4699-4708 (1930); Menzies, A. W. C. ¹⁴⁸

6	0.005604 0.01315 0.01315 0.01315 0.039489 0.08956 0.08956 0.08982 0.0605
×	0.005207 0.005207 0.005338 0.037261 0.05337 1.17906 2.28186 2.28186 2.28186 3.3609 9.4487 11.854 11.858 11.
7	0.004835 .009875 .009875 .009875 .0051844 .051848 .170469 .170469 .170469 .170469 .170469 .170469 .170469 .170469 .17049
9	0.004487 0.009218 0.009218 0.058588 0.058588 0.058588 0.058588 0.058588 0.058588 0.058588 0.058588 0.058588 0.05864 0.05858
LC)	0.004162 .008606 .008606 .013222 .055445 .031222 .15531 .24779 .38043 .57047 .5
4	0.008019 0.029499 0.029499 0.02452 0.029409 0.02452 0.028770 0.029409 0.38479 0.38479 0.38479 0.3849 0.3849 0.3849 0.384
3	0.007473 0.014759 0.014759 0.014759 0.0149599 0.085222 1.049599 1.22553 1.2255
2	0.006960 0.006960 0.016821 0.046881 0.046881 0.046881 0.046881 0.046881 0.046881 0.048831 0.048831 0.04888 0.0
1	0.006479 0.012936 0.012936 0.0142938 0.0442938 0.076718 0.076718 0.076718 1.0362 1.466 3.664 6.254 6.254 6.254 6.254 6.2561 11.662 11.6
0	0.0060273 .0012102 .0012102 .0012103 .1012104 .1012104 .1012104 .1010100 .467350 .46736 .46736 .601923 .10190964 .46736 .601923 .10190964 .1019096
ţ,	11 +

Table 243.—Vapor-pressure of Water (mm-Hg): 0 to −16 °C

From compilation by E. W. Washburn. (See also Tables 242 and 244.)

The following values have been computed by E. W. Washburn ¹⁶⁴ from the values derived by himself for the vapor pressure of ice at the corresponding temperatures (p. 598 and Table 264), the relation between the two being taken as that indicated by the formula

$$\log_{10}\left(\frac{p_w}{p_i}\right) = -\frac{1.1489t}{273.1+t} - 1.330(10^{-5}t^2) + 9.084(10^{-8}t^3)$$

which is based upon the following values:

Specific heat of water = $(1.0092 - 0.001080t + 0.000036t^2)$ cal₁₅ per g·°C Specific heat of ice $^{\alpha}$ = (0.5052 + 0.001861t) cal₁₅ per g·°C

Latent heat of fusion of 18.015 g of ice at 0° C = 1435.5 cal₁₅ (= 333.48 j) Gas-constant, R = 1.9869 cal₁₅ per g-mole·°C = 8.315_2 j/g-mole·°C

The values so obtained for the vapor pressure exceed those given by L. Holborn, K. Scheel, and F. Henning, 166 and based upon work done at the Physikalisch-Technischen Reichsanstalt, by amounts ranging from 0 at 0 °C to 0.008 mm-Hg at -15.9 °C.

When the vapor is mixed with such an amount of atmospheric air that the total pressure is one atm, the vapor pressure exceeds that here given by a small amount, Δp , defined by the equation $\Delta p/p = 0.024/(273 + t)$.* The numbers in this table refer to pure water in contact with its pure vapor. At the end of each line is given, under D, the corresponding average increase in pressure per 0.1 °C decrease in temperature.

Unit of P=1 mm-Hg; of D=0.001 mm-Hg. Temp. = (t_1+t_2) °C -0.0-0.1-0.2-0.3-0.9 $t_{2} \rightarrow$ -0.4-0.5-0.6-0.7-0.8D t_1 4.579 4.546 4.513 4.448 4.289 -04.480 4.416 4.385 4.353 4.320 -32.14.135 -14.258 4.227 4.196 4.165 4.105 4.075 4.045 4.016 3.986 -30.2-23.956 3.927 3.898 3.871 3.785 3.757 3.730 3.702 -28.33.841 3.813 $-\frac{1}{3}$ 3.620 3.647 3.673 3.593 3.567 3.5403.514 3.487 3.461 3.436 -26.33.384 3.359 3.334 3.309 3.284 3.259 3.235 3.410 3.211 3.187 -24.7-5 3.069 3.163 3.139 3.115 3.092 3.046 3.022 3.000 2.976 2.955 -23.2-6 -7 -8 -9 2.887 2.843 2.822 2.931 2.909 2.866 2.800 2.778 2.757 2.736 -21.62.715 2.695 2.674 2.654 2.633 2.613 2.593 2.572 2.553 2.533 -20.12.399 2.514 2.495 2.475 2.380 2.362 2.343 2.456 2.437 2.418 -18.82.326 2.307 2.289 2.271 2.254 2.236 2.219 2.201 2.167 2.184 -17.7-102.149 2.134 2.116 2.099 2.084 2.067 2.050 2.034 2.018 2.001 -16.21.924 -111.987 1.971 1.955 1.939 1.909 1.893 1.878 1.863 1.848 -15.3-121.834 1.819 1.790 1.776 1.734 1.804 1.761 1.748 1.720 1.705 -14.3-131.691 1.678 1.665 1.651 1.637 1.624 1.611 1.599 1.585 1.572 -13.1-141.560 1.547 1.534 1.522 1.511 1.497 1.472 1.485 1.460 1.449 -12.41.425 1.414 1.402 1.390 1.379 1.368 1.356 1.345 1.334 -11.3

In the Weather Review the number 0.5052 was incorrectly printed as 0.5952.

^{*}This coefficient (0.024) differs from that (0.020) given by Washburn 105, 104; the latter is incorrect.

^{151 (}f. W. Thomson (Lord Kelvin), Phil. Mag. (4), 42, 448-452 (1871).

¹⁵² v. Helmholtz, R., Ann. d. Phys. (Wied.), 27, 508-543 (1886).

¹⁵³ Wilson, C. T. R., Phil. Trans. (A), 189, 265-307 (1897).

effect of the pressure that the gas exerts, and (2) the effect of the gas that is dissolved in the liquid. The first is an increase, and the second a decrease, in the pressure of the vapor.

If p_g = partial pressure of the inert gas at the surface of the liquid, p_0 = vapor-pressure of the liquid in the absence of the inert gas, Δp =

Table 244.—Vapor-pressure of Water (millibars): 0 to -50 °C

Adapted from L. P. Harrison. 166 (See also Table 243.)

Computed by means of the formulas (p. 598 and Table 243) given by E. W. Washburn, 164 and for each 0.5 °C.

	Unit o	of $P=1$	millibar =	1000 dy	ne/cm² =	0.75 mm-	Hg. Tem	$\mathbf{p}_{\cdot} = (t_1 +$	$t_2)$ °C	
$t_2 \rightarrow t_1$	0	-1	-2	-3	-4 P	-5	-6	-7	-8	-9
0	6.105	5.677	5.275	4.898	4.546	4.217	3.909	3.620	3.352	3.100
-10 -20	2.866 1.257	2.648 1.153	2.444 1.056	2.255 0.967	2.079 0.885	1.915 0.809	1.763 0.739	1.622 0.674	1.490 0.615	1.370 0.560
-30 -40	$0.510 \\ 0.189$	$0.464 \\ 0.170$	$0.421 \\ 0.153$	0.383 0.138	0.347 0.124	0.314 0.111	0.285 0.0994	0.257 0.0890	0.233 0.0795	0.210 0.0710
-50	0.0633	0.270	0.100	000	0.127	V.111	0.0771	0.0070	0.0770	0.07 20

Table 245.—Vapor-pressure of Water (mm-Hg): 0 to 102 °C (Values in Table 242 are to be preferred.)

From compilation by E. W. Washburn. 167 Cf. also Table 242.

These values are based upon observations made at the Physikalisch-Technischen Reichsanstalt, which were reduced to the temperature scale a of the Reichsanstalt in 1919, and interpolated, as below, by L. Holborn, K. Scheel, and F. Henning. 165 Reproduction is by permission of Vieweg & Sohn.

When the vapor is mixed with such an amount of atmospheric air that the total pressure is 1 atm, the vapor pressure exceeds that here given by the amount Δp , defined as follows: $\Delta p/p = 0.000775 - 3.13(10^{-6}t)$ if $0 \ge$ $t \ge 40 \,^{\circ}\text{C}$; $\Delta p/p = 0.000652 - 8.75(10^{-7}p)$ if $50 \ge t < 90 \,^{\circ}\text{C}$, p being expressed in mm-Hg. The numbers in this table refer to pure water in contact with its pure vapor. At the end of each line is given, under D, the corresponding average increase in the pressure on going from one tabulated value to the next; and under δ is given the amount by which the entry in the zero column exceeds the corresponding value in Table 242, cf. Table 246.

At the triple point corresponding to equilibrium between water, watervapor, and ice-I, t = 0.0099 °C, p = 4.5867 mm-Hg.¹⁶⁸

^e Defined by the melting point of ice (0°), the boiling point of water (100°), and the boiling point of sulphur (taken as 444.55°), all under the pressure of one normal atmosphere; interpolation between these points being made by means of a resistance-thermometer of platinum.¹⁶⁵, p. ⁷

¹⁵⁸a Gudris, N., and Kulikowa, L., Z. Physik, 25, 121-132 (1924).

Hulett, G. A., Z. physik. Chem., 42, 353-368 (1902-1903).
 McHaffie, I. R., and Lenher, S., J. Chem. Soc. (London), 127, 1559-1572 (1925).

	9	ï	+	+11	+17	+23	+27	+32
	D	34.7 36.8 39.1 41.6 44.2	50.0 53.2 56.4 50.0	63.5 67.4 71.3 75.6 80.1	84.6 89.6 94.7 100.0 105.8	111.5 117.7 124.1 130.9 137.9	145.3 153.0 161.0 169.4 178.1	187.1 196.8 206.6 216.9 227.7
	6.0	4.890 5.256 5.645 6.058 6.498	6.965 7.462 7.990 8.551 9.147	9.779 10.449 11.158 11.910 12.706	13.547 14.438 15.380 16.374 17.427	18.536 19.707 20.941 22.243 23.616	25.060 26.582 28.185 29.870 31.642	33.503 35.462 37.518 39.677 41.942
$(t_1 + t_2)$ °C	9.0	4.855 5.219 5.605 6.015 6.453	6.917 7.411 7.936 8.494 9.086	9.714 10.380 11.085 11.833 12.624	13.461 14.347 15.284 16.272 17.319	18.422 19.587 20.815 22.110 23.476	24.912 26.426 28.021 29.697 31.461	33.312 35.261 37.308 39.457 41.710
P. Temp. =	0.7	4.820 5.181 5.565 5.973 6.408	6.869 7.360 7.882 8.437 9.025	9.649 10.312 11.013 11.756 12.543	13.375 14.256 15.188 16.171 17.212	18.309 19.468 20.690 21.977 23.337	24.764 26.271 27.858 29.525 31.281	33.122 35.062 37.099 39.237 41.480
ated digit of	9.0	4.785 5.144 5.525 5.931 6.363	6.822 7.309 7.828 8.380 8.965	9.585 10.244 10.941 11.680 12.462	13.290 14.166 15.092 16.071 17.105	18.197 19.349 20.565 21.845 23.198	24.617 26.117 27.696 29.354 31.102	32.934 34.864 36.891 39.018 41.251
in last tabulated	0.5	5.107 5.107 5.486 5.889 6.318	6.775 7.259 7.775 8.323 8.905	9.521 10.176 10.870 11.604 12.382	13.205 14.076 14.997 15.971 16.999	18.085 19.231 20.440 21.714 23.060	24.471 25.964 27.535 29.184 30.923	32.747 34.667 36.683 38.801 41 023
D and $\delta = 1$	0.4 P	5.070 5.070 5.447 5.848 6.274	6.728 7.209 7.722 8.267 8.845	9.458 10.109 10.799 11.528 12.302	13.121 13.987 14.903 15.871 16.894	17.974 19.113 20.316 21.583 22.922	24.326 25.812 27.374 29.015 30.745	32.561 34.471 36.477 38.584 40.796
l mm-Hg; of	0.3	4.681 5.034 5.408 5.807 6.230	6.681 7.160 7.669 8.211 8.786	9.395 10.042 10.728 11.453 12.223	13.037 13.898 14.809 15.772 16.789	17.863 18.996 20.193 21.453 22.785	24.182 25.660 27.214 28.847 30.568	32.376 34.276 36.272 38.369 40.569
Unit of P =	0.2	4.647 4.998 5.370 5.766 6.187	6.635 7.111 7.617 8.155 8.727	9.333 9.976 10.658 11.379 12.144	12.953 13.809 14.715 15.673 16.685	17.753 18.880 20.070 21.324 22.648	24.039 25.509 27.055 28.680 30.392	32.191 34.082 36.068 38.155 40.344
	0.1	4.613 4.962 5.332 5.725 6.144	6.589 7.062 7.565 8.100 8.668	9.271 9.910 10.588 11.305 12.065	12.870 13.721 14.622 15.575 16.581	17.644 18.765 19.948 21.196 22.512	23.897 25.359 26.897 28.514 30.217	32.007 33.888 35.865 37.942 40.121
	0.0	4.579 4.926 5.294 5.685 6.101	6.543 7.013 7.513 8.045 8.609	9.209 9.844 10.518 11.231 11.987	12.788 13.634 14.530 15.477 16.477	17.535 18.650 19.827 21.068 22.377	23.756 25.209 26.739 28.349 30.043	31.824 33.695 35.663 37.729 39.898
) ئىرىي	011264	20/80	011224	112 113 113 113	82222	28,278,23	332333

28 888 82	T-822824	828824	4444	44444	38 37 35 38 37 35
633.90 657.62 682.07 707.27 733.24 760.00	525.76 546.05 566.99 588.60 610.90	92.51 149.38 233.7 355.1	71.88 75.65 79.60 83.71 88.02	55.324 58.34 61.50 64.80 68.26	42.175 44.563 47.067 49.692 52.442
636.24 660.03 684.55 709.83 735.88 762.72 790.37	527.76 548.11 569.12 590.80 613.17	97.20 156.43 243.9 369.7	72.25 76.04 80.00 84.13 88.46	55.61 58.65 61.82 65.14 68.61	42.409 44.808 47.324 49.961 52.725
638.59 662.45 687.04 712.40 738.53 765.45 793.18	5.29.77 550.18 571.26 593.00 615.44	102.09 163.77 254.6 384.9	72.62 76.43 80.41 84.56 88.90	55.91 58.96 62.14 65.48 68.97	42.644 45.054 47.582 50.231 53.009
640.94 664.88 689.54 714.98 741.18 768.19 796.00	531.78 552.26 573.40 595.21 617.72	107.20 171.38 265.7 400.6	72.99 76.82 80.82 84.99 89.34	56.21 59.27 62.47 65.82 69.93	42.880 45.301 47.841 50.502 53.294
643.30 667.31 692.05 717.56 743.85 770.93 798.82	533.80 554.35 575.55 597.43 620.01	112.51 179.31 277.2 416.8	73.36 77.21 81.23 85.42 89.79	56.51 59.58 62.80 66.16 69.69	43.117 45.549 48.102 50.774 53.580
645.67 669.75 694.57 720.15 746.52 773.68 801.66	P 0.5 535.82 556.44 577.71 599.66 622.31	P 118.04 187.54 289.1 433.6	73.74 77.60 81.64 85.85 90.24	56.81 59.90 63.13 66.51 70.05	43.355 45.799 48.364 51.048 53.867
648.05 672.20 697.10 722.75 749.20 776.44 804.50	537.86 558.53 579.87 601.89 624.61	123.80 196.09 301.4 450.9	74.12 78.00 82.05 86.28 90.69	57.11 60.22 63.46 66.86 70.41	43.595 46.050 48.627 51.323 54.156
650.43 674.66 699.63 725.36 751.89 779.22 807.35	539.90 560.64 582.04 604.13 626.92	7 129.82 204.96 314.1 468.7	74.50 78.40 82.46 86.71 91.14	57.41 60.54 63.79 67.21 70.77	43.836 46.302 48.891 51.600 54.446
652.82 677.12 702.17 727.98 754.58 782.00 810.21	541.95 562.75 584.22 606.38 629.24	136.08 214.17 327.3 487.1	74.88 78.80 82.87 87.14 91.59	57.72 60.86 64.12 67.56 71.14	44.078 46.556 49.157 51.879 54.737
655.22 679.59 704.71 730.61 757.29 784.78 813.08	544.00 564.87 586.41 608.64 631.57	142.60 223.73 341.0 506.1	75.26 79.20 83.29 87.58 92.05	58.03 61.18 64.46 67.91 71.51	44.320 46.811 49.424 52.160 55.030
237.2 244.5 252.0 259.7 267.6 275.7 283.9	202.9 209.4 216.1 223.0 230.0	568.7 843.2 121.4 170.7	37.7 39.5 41.1 43.1 44.6	30.2 31.6 33.0 34.6 36.2	238.8 250.4 262.5 275.0 288.2
-10 +6	-10	1 1 2 8	+	+36	+35

amount by which the equilibrial pressure of the vapor has been increased by the presence of the inert gas, $\rho =$ density of the liquid, $T \circ K =$ absolute temperature, R = the gas-constant expressed in appropriate units, M = molecular weight of the vapor, and n_g and $n_l =$ number of molecules of gas and of liquid, respectively, that make up the liquid phase, then if Δp is small, it is quite closely given by the formula $\Delta p/p_0 = Mp_g/\rho RT - n_g/(n_g + n_l)$. A more accurate formula is $\log_e[(n_g + n_l)p'/n_lp_0] = Mp_g/\rho RT$, where $p' = p_0 + \Delta p$, and ρ' is the density of the liquid at a pressure midway between p_0 and $(p' + p_g)$. It must be remembered that p_g is the actual partial pressure at the surface of the liquid, and that n_g is the actual number of molecules dissolved in the liquid; if the liquid is boiling, these values are quite different from what they would be under other conditions. Under suitable boiling conditions they approach, and may become, zero.

Apparently no direct determination of the equilibrial pressure of the

Table 246.—Comparison of Smoothed Values for the Vapor-pressure of Water (mm-Hg, atm)

In this table several sets of smoothed values for the vapor-pressure (P) of water are compared with those preferred ones $(P_{\rm OM})$ defined by formula (1) of Osborne and Meyers, and given in Table 242. Some of these sets represent values computed by means of smoothing equations that have been published; all have been taken from published tables of adjusted values. None of them are directly observed values. Graphical comparisons of the observed values have been published by N. S. Osborne, 160 and by N. S. Osborne and C. H. Meyers. 156

 $P=P_{\rm OM}+\delta$, where $P_{\rm OM}$ is the value defined by the Osborne-Meyers formula. When P has been published with less than 5 significant figures (the number here given for $P_{\rm OM}$) or in such units that the last specified digit does not suffice to determine the fifth digit when P is expressed in the unit here used, then the surplus or undetermined digits in δ are depressed; when the significant figures in δ are all depressed, they are preceded by a 0 in the normal position. For example, the KS value for $160~{\rm ^{\circ}C}$ is $6.0996+0_4=6.100$ atm.

Unit of p = 1 mm-Hg; of P - 1 atm = 1.01325 bars = 1.03323 kg*/cm²; of $\delta = 1$ unit in last place in $b_{\rm DM}$ or $P_{\rm DM}$

				prace in	POM O	or FOM				
Refe	erencesa→	ICT ₁	IST	SKG_2	Ref	erences ^a →	ICT ₁	IST	SKG_2	\mathbf{SM}
ŧ	POM		δ		ŧ	p_{OM}		(5	
0	4.5807	-1_{7}	$+0_{4}$	$+8_{6}$	50	92.490	$+2_{0}$	$+2_{0}$	+13	-22_{0}
5	6.5360	+70		$+11_{9}$	55	118.04	0		0	-17
10	9.1975	$+11_{5}$	+6s	$+14_{0}$	60	149.39	-1	+2	-1	-26
15	12.771	+17		+18	65	187.56	-2		-2	-37
20	17.512	+23	+16	+22	70	233.72	-0_{2}	0	-3	-28
25	23.729	+27		+24	75	289.13	-0_{3}	0	-3	-35
30	31.792	+32	+24	+25	80	355.22	— 1 ₂	0	-4	-32
35	42.139	+36		+26	85	433.56	$+0_4$		-5	-2
40	55.288	+ 36	+29	+23	90	525.86	-10	0	-4	+8
45	71.840	$+4_{o}$		+20	95	634.00	-10		-3	

¹⁵⁶ Osborne, N. S., and Meyers, C. H., J. Res. Nat. Bur. Stand., 13, 1-20 (RP691) (1934) \rightarrow Mech. Eng., 56, 207-209 (1934).

Table 246.—(Continued)

					,	/			
Refe	rences ^a →	OSFG	SKG	KS	1CT ₂	JF	WT	EC	С
t	P_{OM} —				δ-				
100	1.0000	0	0		0		0		
110	1.4139	-1	ŏ		ŏ		ő		
120	1.9594	-1 -1	+1		ŏ		+1		
130	2.6658	0	+4		+2		+4		
140	3.5663	+1	+5		$+\tilde{0}_{7}$		$+0_{7}$		
150	4.6975	+ 2	+8		+05		+ 05		
160	6.0996	44		+04	+ 0 ₅		+05		
170	7.8167	+4	$^{+11}_{+13}$	$+2_{3}$	$+0_{3}$		$+1_{3}$	+79	
180	9.8960	+2	+13 + 14	-3_{0}	-1_{0}	$+0_{50}$	00	+126	
190	12.388	Ť 0	+2	-10°	$-\frac{10}{-2}$	00	-1^{00}	+12	
				-12		-			1 12
200	15.347	0	+2		-6	-0_{7}	-4	+8	$+13_{5}$
210 220	18.830 22.898	-1^{0}	+1	$^{+12}_{-8}$	−7 −9	— 0 ₆	$-4 \\ -5$	+5 +5	$^{+17_8}_{+22_3}$
230	27.613	$-1 \\ -1$	0	-23	-10	— 0. — 1.	$-5 \\ -5$	+8	$+26_7$
240	33.042	-1	-1^{0}	-35	-15	-1_{0}^{10}	_9	+6	$+31_{1}$
250	39.256	0	-2	-25	-22	-2_{0}	-15	-4	+354
260	46.326	+ 2	-3	-15	-26	-2.	-17	-1	$+38_{5}$
270	54.331	+3	-4	$-20 \\ -41$	-40	$-3_{\rm s}$	$-28 \\ -43$	+14	$+39_{3}$
280	63.352	$+\frac{1}{2}$	-6	-41	-57	-5_{5}		$^{+14}_{-4}$	$+37_{3}$
290	73.475	-3	-9		-5_{5}	-5_5	-3_{5}		$+32_{6}$
300	84.793	-5	-14	+4	-1_{3}	-0_{1}	$+0_{7}$	-11	+17s
310	97.406	-4	-21	+30	$-0_{\rm o}$	$+0_{7}$	+2.	+8	+87
320	111.42	0	-3	+5	+1		+4	+2	-1_{7}
330	126.96	0	-3	+10	+3		+7	$-\frac{1}{2}$	-4_{e}
340	144.17	0	-4	+16	+3		+7	-2	-7 ₉
350	163.20	0	-3	+16	-4		+1	+2	-9_{6}
360	184.29	0	-2	+15	-22		-16	+1	-9_{5}
370	207.77	+1	-3	+7	-28		-21	+7	$+1_{5}$
371	210.27	+1	_	+9	-29		-21	+9	
372	212.80	+1	-5	+12	+29		-21	+8	
373	215.37	0		+15	+28		-20	+8	
374	217.98	-2	-10	+ 20	+26		-18	+7	+13s

a References:

Callendar, H. L., Proc. Inst. Mech. Eng., 1929, 507-527 (1929).
Egerton, A., and Callendar, G. S., Phil. Trans. (A), 231, 147-205 (A698) (1932).
From compilation by Washburn, E. W. 197 WT; see Table 245.
From the compilation by Keyes, F. G., Int. Crit. Tables, 3, 233 (1928), and derived by him from WT by the application of corrections required to make the temperature scale correspond to that on which the boiling point of sulphur under a pressure of 1 atm is 444.60 °C instead of 444.55 °C, which is assumed in WT.
Int. Skeleton Steam Tables, 1934, see Table 260.
Jakob, M., and Fritz, W., Techn. Mech. u. Thermodyn., 1, 173-183, 236-240 (1930); Forsch. Gebiete Ingenicurus, 4, 295-299 (1933).
Keyes, F. G., and Smith, L. B., Mech. Eng., 52, 124-127 (1930).
Osborne, N. S., Stimson, H. F., Fiock, E. F., and Ginnings, D. C., Bur. Stand. J. Res., 10, 155-188 (RP523) (1933).
Smith, L. B., Keyes, F. G., and Gerry, H. T., Proc. Amer. Acad. Arts. Sci., 69, 137-168 (1934); they publish two smoothing equations—(1) for t > 100 °C and (2) for t < 100 °C, as here indicated by subscripts—and extended tables of values of p and of dp/dt.
Smith, A., and Menzies, A. W. C., Ann. d. Phys. (4), 33, 971-978 (1910). IST JF KS OSEG SKG Smith, A., and Menzies, A. W. C., Ann. d. Phys. (4), 33, 971-978 (1910). Holborn, L., Scheel, K., and Henning, F.166

vapor of water in contact with an inert gas is available, but insofar as it is allowable to regard water-vapor as an ideal gas the increase Δp may be computed by means of the formula $\Delta p/p_0 = (\rho' - \rho_0)/\rho_0$, where p_0 is the value of the vapor-pressure of water when no inert gas is present, ρ_0 is the

^b Above t = 100 °C the IST values are identical with P_{OM} .

corresponding density of the pure saturated vapor, and ρ' is the observed density of water-vapor saturated in the presence of the inert gas; all corresponding to the same temperature. These values of $\Delta p/p_0$ may be read directly from the values of ρ'/p_0 given in Table 252. It will be noticed that the several sets of comparable values there given are discordant both with one another and with the values computed by means of the preceding formula.

Curvature of Surface, Effect of.—If p = equilibrial pressure of thevapor in contact with a spherical surface of the liquid (radius = r), p_0 = that of the vapor in contact with a flat surface, $\gamma = \text{surface-tension}$ of the liquid, and the other symbols have the significance already stated, then $(p-p_0)/p_0 = \pm \left[\gamma \rho_0/(\rho-\rho_0)p_0\right](1/r_1+1/r_2) = \pm 2\gamma/\rho RTr$ approximately, if $(p - p_0)$ and $(p - p_0)$ are small, and $r_1 = r_2 = r$, r_1 and r_2 being the principal radii of curvature of the surface; the plus sign is to be taken if the concavity of the surface is toward the liquid. 151 (If r is so small that $p - p_0$ is not small in comparison with p_0 , then the more exact expression $\log_e(p/p_0) = (\gamma/RT_P)(1/r_1 + 1/r_2)$ must be used; see R. v. Helmholtz ¹⁵² or C. T. R. Wilson. ¹⁵³) Obviously, the unit of mass appearing in R must be the same as that in ρ . For water at 20 °C $(T = 293.1 \text{ °K}), \gamma = 72.75 \text{ dynes/cm}, \rho = 0.9984 \text{ g/cm}^3, R = 8.315 \times 10^7$ erg/g-mole.°C = 4.615×10^6 erg/g.°C, whence $r(p - p_0)/p_0 = 0.001077 \mu$ if the concavity is turned away from the liquid $(1 \mu = 0.001)$ mm). Whence the following:

Certain observations by W. A. Patrick and his associates indicate that over concave menisci in very small tubes (r = a few microns) the value of $(p_0 - p)$ exceeds that demanded by this formula. They are inconsistent with the observations of N. Gudris and L. Kulikowa, 153a and seem to be

Table 247.—Thermal Rate of Variation in the Vapor-pressure of Water (atm) 156

These values, computed by means of formula (2), are believed to be the most accurate available; they supersede the slightly different ones previously published by Osborne, Stimson, Fiock, and Ginnings.¹⁷⁰ In Table 248 certain of these values are expressed in mm-Hg, and other sets of values are compared with them.

The authors give values for each °C in kg*/cm² as well as in atm, and for each °F in lb*/in². And Osborne, Stimson, and Ginnings ¹⁷¹ give the values of T(dP/dT) in Int. joules/cm³ for these same values of dP/dT (steps not greater than 5 °C; range, 100 to 374.15 °C).

¹⁵⁷ Adamson, A., Mem. Proc. Manchester Lit. Phil. Soc., 76, 1-9 (1931).

¹⁵⁸ Batschinski, A., Nature, 119, 198 (1927).

¹⁵⁰ Fischer, V., Z. Physik, 43, 131-151 (1927).

¹⁰⁰ Hofbauer, P. H., Atti pontif. Acc. sci. nuovi Lincei, 84, 353-363, 581-586 (1931).

¹⁶¹ Kiréeff, V., Jour. de Phys. (7), 3, 150D (A) (1932) ← Jour. chim. phys. Russe, 1, 241-248 (1930).

	9	0.4099 .7636 1.3514 2.2841 3.7043 5.788	8.747 12.825 18.297 25.465 34.655	46.210 60.49 77.84 98.66 123.28	152.08 185.40 223.60 267.00 315.92	370.69 431.61 498.98 573.2 654.4	743.2 839.8 944.7 1058.6 1181.9	1315.6 1460.6 1618.3 1790.8 1981.7	2197.7 2455.3
bộ	8	0.3840 .7193 1.2792 2.1715 3.5356 5.544	8.405 12.359 17.679 24.663 33.636	44.938 58.93 75.96 96.41 120.64	149.00 181.86 219.55 262.42 310.77	364.94 425.23 491.95 565.4 646.0	733.9 829.8 933.8 1046.8 1169.1	1301.7 1445.5 1601.9 1772.8 1961.6	2174. 6 2426.6
760000 mm-H	7	0.3595 .6772 1.2102 2.0636 3.3733 5.309	8.074 11.907 17.077 23.881 32.639	43.693 57.40 74.11 94.20 118.03	145.97 178.36 215.56 257.89 305.68	359.25 418.92 484.98 557.8 637.6	724.8 819.8 923.0 1035.0	1288.0 1430.6 1585.6 1755.0	2151.9 2398.7
1 mılliatın = 1.0332 g*/cm² = 1.0136 millibar = 0.760000 mm·Hg. Temp. = $(t_1 + t_2)$ °C (Int. scale)	9	0.3364 .6372 1.1445 1.9602 3.2172 5.082	7.754 11.468 16.492 23.118 31.666	42.474 55.90 72.29 92.03 115.47	142.98 174.91 211.61 253.41 300.65	353.63 412.67 478.07 550.2 629.3	715.7 809.9 912.3 1023.4 1143.9	1274.3 1415.8 1569.5 1737.3	2129.6 2371.6
$^*/\text{cm}^2 = 1.013$ (Int. scale)	dP/dt 5	0.3146 .5993 1.0818 1.8612 3.0672 4.8624	7.444 11.043 15.922 22.374 30.715	41.282 54.43 70.51 89.89 112.95	140.04 171.51 207.72 248.99 295.67	348.06 406.48 471.24 542.6 621.0	706.7 800.2 901.7 1011.9 1131.4	1260.8 1401.1 1553.5 1719.8 1902.8	2107.6 2345.2
m = 1.0332 g $(t_1 + t_2)$ °C (4 dP	0.5633 1.0220 1.7665 2.9230 4.6510	7.144 10.630 15.368 21.649 29.786	40.115 52.99 68.76 87.80 110.74	137.13 168.15 203.87 244.63 290.75	342.56 400.36 464.47 535.2 612.9	697.8 790.5 891.2 1000.5 1119.0	1247.4 1386.6 1537.7 1702.5 1883.6	2085.9 2319.4 2630.7
	6	0.5292 .9651 1.6758 2.7846 4.4472	6.854 10.229 14.830 20.943 28.880	38.974 51.57 67.04 85.74 108.03	134.28 164.85 200.08 240.32 285.89	337.11 394.30 457.77 527.8 604.8	689.0 780.8 880.7 989.2 1106.7	1234.1 1372.1 1522.0 1685.3 1864.6	2064.5 2294.1 2584.1
Unit of $dP/dt = 1$ milliatm/°C.	2	0.4968 .9109 1.5891 2.6516 4.2509	6.574 9.841 14.307 20.255 27.995	37.858 50.19 65.35 83.71 105.62	131.46 161.58 196.34 236.06 281.08	331.73 388.30 451.13 520.5 596.8	680.2 771.3 870.4 977.9 1094.5	1220.9 1357.8 1506.4 1668.3 1845.9	2043.4 2269.3 2548.6
Unit of dP/d	1	0.4662 .8593 1.5063 2.5240 4.0618	6.303 9.465 13.799 19.585 27.131	36.766 48.836 63.70 81.72 103.26	128.69 158.37 192.64 231.85 276.33	326.40 382.37 444.56 513.3 588.8	671.6 761.8 860.1 966.8 1082.4	1207.8 1343.6 1491.0 1651.5 1827.3	2022.5 2245.0 2515.9
	0	0.4373 .8103 1.4207 2.4015 3.8796	6.041 9.100 13.305 18.932 26.288	35.699 47.509 62.08 79.76 100.94	125.96 155.20 189.00 227.70 271.64	321.13 376.50 438.05 506.1 581.0	662.9 752.5 849.9 955.7 1070.4	1194.8 1329.5 1475.7 1634.8 1809.0	2002.0 2221.2 2485.0
	ţ,	+ 10 20 30 40 40	2000 2000 8000 8000	120 130 140	150 160 170 180	220 220 230 240	250 270 280 280	300 310 320 330 340	350 360 370

inconsistent with the results obtained in another field by R. Buckley (see discussion on pp. 513, 518).

Tension, Effect of.—G. A. Hulett 154 has reported observations which he

Table 248.—Comparison of Thermal Rates of Variation in the Vapor-pressure of Water (mm-Hg)

 $dp/dt = (dp/dt)_{\rm OM} + \delta$ where $(dp/dt)_{\rm OM}$ is the corresponding value from Table 247.

1 mm-Hg = 1.3158 milliatm = 1.3564 g*/cm² = 1.3332 millibars.

	τ	Jnit of (dp/dt) = 1	mm-Hg	$s/^{\circ}C$; of $\delta = 3$	1 unit i	in last place	in (dp/dt) _{OM}	
t	$Ref^{a} \rightarrow (dp/dt)$		WT Jŀ		OSFG		$ef^{a} \rightarrow (dp/dt)_{OM}$	JF	J OSFG	_
0 10 20 30 40	0.332 .615 1.080 1.825 2.948	8 +7 +5 0	+ 12 + 20 + 6 + 1 - 1	-1 -7		200 210 220 230 240	244.0 286.1 332.9 384.6 441.6	0 -0 0 -1 -6	-2 0 0 0 0 0 0 0	
50 60 70 80 90	4.591 6.916 10.11 14.38 19.98	$ \begin{array}{r} -1 \\ -2 \\ 0 \\ -1 \\ 0 \end{array} $	-4 -8 -1 0 -1	$ \begin{array}{r} -5 \\ -10 \\ -3 \\ 0 \\ +2 \end{array} $		250 260 270 280 290	503.8 571.9 645.9 726.3 813.5	$ \begin{array}{r} -2 \\ -8 \\ -9 \\ -10 \\ +5 \end{array} $	+1 +1 +1 -3 -2	
100 110 120 130 140	27.13 36.11 47.18 60.62 76.71	$ \begin{array}{c} +1 \\ +1 \\ +1 \\ +1 \\ 0 \end{array} $	$^{+1}_{+1}_{+1}_{0}_{0}$	+3 0 0 +2 -7	$ \begin{array}{c} -1 \\ -1 \\ 0 \\ +1 \\ +1 \end{array} $	300 310 320 330 340	908.0 1010.4 1121.5 1242.4 1374.8	+40 -5	-1 +2 + ; +1 3	
150 160 170 180 190	95.73 118.0 143.6 173.0 206.4	-2	$-3 \\ -5$	$\begin{array}{c} +3 \\ 0 \\ +1 \\ 0 \\ -2 \\ 1 \end{array}$	+1 0 0 0 0	350 360 365 370 374	1521.5 1688.1 1782.4 1888.6 1999.3		3 +6 +11 -3 -181	
ł	I. Mose				44.43.	_				
	96.5 97 97.5	_{ф/dt) ом} 24.434 24.806 25.182	$ \begin{array}{r} \delta \\ +55 \\ +13 \\ -20 \end{array} $	98 98.5 99	(dp/dt) _{OM} 25.563 25.948 26.338	δ 4 5 5	5 99.5 4 100	(dp/dt), 26.73 27.13 27.53	32 - 38 31 - 9	
а	 References: Jakoh, M., ForschArb. Gebiete Ingenieurw., 310, 9-19 (1928). Jer Jakob, M., and Fritz, W., Techn. Mech. Thermodynam., 1, 173-183, 236-240 (1930). OSFG Osborne, N. S., Stimson, H. F., Fiock, E. F., and Ginnings, D. C., Bur. Stand. J. Res., 10, 155-188 (RP523) (1933). SKG Smith, L. B., Keyes, F. G., and Gerry, H. T., Proc. Amer. Acad. Arts Sci., 69, 137-168 (1934). WT Holborn, L., Scheel, K., and Henning, F., "Warmetabellen," 1919. 									

Table 249.—Temperature of Saturated Water-vapor: 0.0075 to 225 kg*/cm²

Adapted from a table by N. S. Osborne and C. H. Meyers ¹⁵⁶ and on the same basis as the data in Table 242. The authors give a similar table in terms of $1b*/in^2$ and °F, but do not give the Δ 's. (See also Table 253.)

100 Smith, L. B., Keyes, F. G., and Gerry, H. T., Proc. Amer. Acad. Arts Sci., 69, 137-168 (1934); 70, 319-364 (1935).

Table 249.—(Continued)

Values of t for intermediate values of P may be computed by means of the well-known formula:

$$f(x+h) = f(x) + k \left\{ \Delta_1(x) - \frac{1-k}{2} \Delta_2(x) + \frac{(1-k)(2-k)}{3!} \Delta_3(x) - \frac{(1-k)(2-k)(3-k)}{4!} \Delta_4(x) + \cdots \right\}$$

where k = h/s, s being the length of the successive (equal) steps in the table, and Δ_n being the nth difference (p. 656). The first and second differences are tabulated on the same lines as the P's to which they correspond. They will be sufficient except where Δ_2 is varying rapidly; in which case, higher differences should be computed and used.

For example, taking one of the worst cases, find the temperature at which the saturation pressure is 0.007721_3 kg*/cm². The nearest tabular value below the assigned pressure is P = 0.0075, corresponding to t = 2.590 °C, $1000\Delta_1 = 4117$, $1000\Delta_2 = -828$, $1000\Delta_3 = +292$, $1000\Delta_4 = -135$, $1000\Delta_5 = +77$; $h = 0.000221_3$, s = 0.0025, hence k = 0.0884. Putting these in the formula, one obtains the value

$$t = 2.590 + \frac{0.0884}{1000} \{4117 + 377 + 85 + 29 + 13 + \cdots\} = 2.998_5 \, ^{\circ}\text{C}$$

Whereas it is seen from Table 242 that at $3 \,^{\circ}\text{C}$ P = 0.007473 atm = 0.007721_3 kg*/cm². Hence, the computed value differs from the true by only $0.001_5 \,^{\circ}\text{C}$. Had all differences beyond the second been neglected, the value 2.987 $\,^{\circ}\text{C}$ would have been found.

Similarly for $P=0.018512_4$ kg*/cm² one obtains t=15.997 °C if only the first and second differences are used; 16.000 °C if the third is included. From Table 242 it is seen that at 16 °C, P=0.017917 atm = 0.018512₄ kg*/cm². Here the computed value is in error by less than 0.001 °C when only 3 differences are used.

Unit of $P =$	1 kg*/cm2	= 0 967841	atm = 735.559	mm-Hg = 0.90665	bar.	Temp. = t °C,	Int. scale
P	ŧ	1000 Δ1	$-1000 \Delta_{2}$	P	ŧ	1000 Δ1	$-1000 \Delta_{2}$ -
0.0075	2.590	4117	828	0.065	37.302	2 1370	83
.0100	6.707	3289	536	.070	38.672	2 1287	72
.0125	9.996	2753	379	.075	39.959		63
.0150	12.749	2374	280	.080	41.17		58
.0175	15.123	2094	220	.085	42.320		51
.0200	17.217	1874	174	.090	43.420		46
.0225	19.091	1700	142	.095	44.46		
.0250	20.791	1558	119	.100	45.46		140
.0275	22.349	1439		.11	47.33		120
.030	23.788	2587	304	.12	49.06		103
.035	26.375	2283	236	.13	50.67		90
.040	28.658	2047	190	.14	52.18		7 8
.045	30.705	1857	154	.15	53.59		<i>7</i> 0
.050	32.562	1703	129	.16	54.93		63
.055	34.265	1574	111	.17	56.20		55
.060	35.839	1463	93	.18	57.41	6 1152	51

Table 249.—(Continued)

P	ŧ	1000 Δ1	-1000 A2	P	ŧ	1000 Δ1	-1000 A ₂
0.19	58.568	1101	46	4.3	145.538	839	15
.20	59.669	1055	41	4.4	146.377	824	15
.21	60.724	1014	39	4.5	147.201	809	14
.22	61.738	975	36	4.6	148.010	795	13
.23	62.713	939	32	4.7	148.805	782	13
.24	63.652	907	31	4.8	149.587	769	13
.25	64.559	876	01	4.9	150.356	756	
.26	65.435	1671	99	5.0	151.112	1477	45
.28	67.106	1572	86	5.2	152.589	1432	41
.30	68.678	1486	77	5.4	154.021	1391	40
.32	70.164	1409	68	5.6	155.412	1351	36
.34	71.573	679	61	5.8	156.763	1315	35
.35	72.252	662	01	6.0	158.078	1280	35 33
.36	72.914	1280	56	6.2	159.358	1247	30
.38	74.194	1224	30	6.4	160.605	1217	29
.40	75.418	2849	256	6.6	161.822	1188	27
.45	78.267	2593	208	6.8	163.010	1161	27
.50	80.860	2385	176	7.0	164.171	1134	24
.55	83.245	2209	149	7.2	165.305	1110	24
.60	85.454	2060	128	7.4	166.415	1086	22
.65	87.514	1932	112	7.6	167.501	1064	22 22
.70	89.446	1820	98	7.8 7.8	168.565	1042	20
.75	91.266	1722	88	8.0	169.607	1042	20
.80	92.988	1634	78	8.2	170.629	1002	18
			71	8.4	171.631	984	18
.85 .90	94.622 96.178	1556 1485	61	8.6	172,615	966	18
.95	97.663	1424	01	8.8	173.581	948	
1.0		2677	194	9.0	174.529	932	16 16
1.1	99.087 101.764	2483	166	9.0	175.461	916	10
1.2	104.247	2317	142	9.4	176.377	901	15 15
1.3			125	9.4	170.377	886	14
1.4	106.564 108.739	2175 2050		9.0 9.8	177.278 178.164		14
1.5		1941	109	10.0	179.036	872 2122	79
1.6	110.789 112.730	1843	98 86	10.5	181.158	2043	72
	114.730				101.100		12
1.7 1.8	114.573	1757 1679	78	11.0	183.201	1971	65
1.6	116.330 118.009	1608	71 65	11.5 12.0	185.172 187.078	1906 1844	62 57
2.0	119.617	1543	58	12.5	188,922	1787	53
2.1	121.160	1485	55	13.0	190.709	1734	49
2.2							49
2.2	122.645	1430	49	13.5	192.443 194.128	1685	
2.3	124.075 125.456	1381 1335	46 44	14.0	194.128	1638	44 40
2.5	126.791	1291	38	14.5 15.0	195.766 197.360	1594 1554	39
2.6	128.082	1253	38	15.5	198,914		37
2.7	129.335	1215	30	15.5	200.429	1515 1478	3/
2.8	130.550	1180	35	16.0 16.5	201.907		34
2.9	131.730	1148	32 31			1444	32 32
3.0	132.878	1117	29	17.0 17.5	203.351 204.763	1412	32
3.1	133.995	1088	29 27	18.0		1380	29 28 27
3.2	135.083	1061	2/	10.0	206.143	1351	28
3.3	136.144	1036	25 24	18.5 19.0	207.494	1323	27 26
3.4	137.180	1012	24		208.817	1296	20
3.5	138.192	988	24 22 21	19.5 20.0	210.113	1270 1246	24
3.6	139.180	966	21	20.5	211.383		23
3.7	140.146		10		212.629	1223	22 22
3.7 3.8		945	19	21.0	213.852	1201	22
3.9	141.091 142.017	926 906	20 17	21.5	215.053	1179	21 19
4.0	142.017	889	17	22.0	216.232	1158	19
4.0	142.923	872	18	22.5 23.0	217.390	1139	19
4.2	144.684	854		23.5	218.529	1120	19
7.2	144.004	034	15	23.3	219.649	1101	17

Table 249.—(Continued)

P	t	1000 Δ1	-1000 A ₂	P		1000 Δ1 -	−1000 ∆₂
24 0	220 750	1084	17	78	291.864	881	8
24 5	221 834	1067	17	79	292 745	873	8
25 0	222 901	1050	15	80	293 618	865	9
25 5	223 951	1035	15	81	294,483	856	8
26.0	224 986	1020	15 15	82	295 339	848	7
26 5	226 006	1005	15	83	296 187	841	8
27.0	227 011	990	13	84	297 028	833	8
27 5	228 001	977	13	85	297 861	825	7
28 0	228 978	964	14	86	298.686	818	7
28.5	229 942	950	12	87	299 504	811	7
29 0	230 892	938	12	88	300 315	804	7
29 5	231 830	926		89	301 119	797	7
30 0	232 756	1817	45	90	301 916	790	9878877777767576
31	234 573	1772	42	91	302 706	783	6
32	236 345	1730	39	92	303 489	777	7
33	238 075	1691	38	93	304 266	770	5
34	239 766	1653	37	94	305 036	765	7
35	241 419	1616	33	95	305 801	758	6
36	243 035	1583	32	96	306 559	752	6
37	244 618	1551	32	97	307 311	746	6
38	246 169	1519	29	98	308 057	740	6
39	247 688	1490	28	99	308 797	734	J
40	249 178	1462	28	100	309 531	1452	22
41	250 640	1434	25	102	310 983	1430	21
42	252 074	1409	25	104	312 413	1409	20
43	253 483	1384	24	106	313 822	1389	20
44	254 867	1360	23	108	315 211	1369	20
45	256 227	1337	22	110	316 580	1349	18
46	257 564	1315	21	112	317 929	1331	18
47	258 879	1294	21	114	319 260	1313	18
48	260 173	1273	19	116	320 573	1295	17
49	261 446	1254	20	118	321 868	1278	16
50	262 700	1234	17	120	323 146°	1262	
51	263 934	1217	19	122	324 408	1245	15
52	265 151	1198	16	124	325 653	1230	15
53	266 349	1182	18	126	326 883	1215	17 15 15 15 15
54	267 531	1164	15	128	328 098	1200	15
55	268 695	1149	16	130	329 298°	1185	14
56	269 844	1133	14	132	330 483	1171	13
57	270 977	1119	16	134	331 654	1158	14
58	272 096	1103	14	136	332 812	1144	13
59	273 199	1089	13	138	333 956	1131	13
60	274 288	1076	14	140	335 087°	1118	12 .
61	275 364	1062	13	142	336 205	1106	13
62	276 426	1049	12	144	337 311	1093	11
63	277 475	1037	14	146	338 404	1082	12
64	278 512	1023	11	148	339 486	1070	12
65	279 535	1012	10	150	340 556°	1058	11
66	280 547	1002	13	152	341 614	1047	11
67	281 549	989	11	154	342 661	1036	10
68	282 538	978	11	156	343 697	1026	11
69	283 516	967	10	158	344 723	1015	11
70	284 483	957	10	160	345 738°	1004	9
71	285 440	947	11	162	346 742	995	11
72	286 387	936	9	164	347 737	984	9
73	287 323	927	10	166	348 721	975	10
74	288 250	917	9	168	349 696	965	9
75	289 167	908	9	170	350 661°	956	10
76	290.075	899	9	172	351 617	946	8
77	290.974	890	9	174	352.563	938	10

				(Communica)			
P	•	$1000 \; \Delta_1$	$-1000 \Delta_{2}$	P	ı	1000 Δ1	-1000 A ₂
17 6	353.501	928	9	202	364.914	823	8
178	354.429	919	8	204	365.737	815	8
180	355.348°	911	9	206	366.552	807	7
182	356.259	902	8	208	367.359	800	8
184	357.161	894	8	210	368.159	792	6
186	358.055	886	9	212	368.951	786	9
188	358.941	877	8	214	369.737	777	6
190	359.818 <i>°</i>	869	8	216	370.514	771	8
192	360.687	861	8	218	371.285	763	8
194	361.548	853	7	220	372.048	755	8
196	362.401	846	9	222	372.803	747	
198	363.247	837	7	224	373.550	370	
200	364.084°	830	7	225	373.920	•. •	

Table 249.—(Continued)

regarded as indicating that the subjecting of water to a hydrostatic tension decreases its vapor-pressure. An effect of that kind is to be expected, but it seems probable that his observations are complicated by the phenomena considered in the preceding paragraphs.

Solute, Effect of.—The partial pressure of water-vapor in equilibrium with an aqueous solution is less than that in equilibrium with pure water at the same temperature (see p. 582). For dilute solutions the reduction in vapor-pressure is approximately proportional to the molecular concentration of the solution.

Adsorbed Water, Effect of.—I. R. McHaffie and S. Lenher 535 have measured the amount of water-vapor adsorbed on glass and on platinum for each of a series of associated temperatures and pressures, and have concluded that the equilibrium pressure is less than the vapor pressure of water unless the adsorbed layer is several hundred molecules thick, if on the glass, and several tens of molecules thick, if on platinum.

Formulas.—Numerous formulas connecting the temperature and the vapor pressure of water have been proposed. The most satisfactory is the one (1) proposed by N. S. Osborne and C. H. Meyers 156 and shown by them to fit the available observations from -5 °C to the critical point.

$$\log_{10}P = A + \frac{B}{T} + \frac{Cx}{T} \left(10^{Dx^2} - 1\right) + E(10^{Fy^{5/4}}) \tag{1}$$

the unit of the saturation pressure (P) being 1 int. atm, the corresponding temperature being t °C (int. scale) and the other quantities having these values: T = t + 273.16, $x = T^2 - K$, y = 374.11 - t, A = + 5.4266514, B = -2005.1, $C = +1.3869(10)^{-4}$, $D = +1.1965(10)^{-11}$, K = +

These values agree with those published by W. Koch ¹⁷⁸ in 1934, but differ from those he published earlier ¹⁷⁴ by amounts ranging from -0.04 to +0.10 °C.

¹⁰⁸ Washburn, E. W., Int. Crit. Tables, 3, 210-212 (1928).

¹⁶⁴ Washburn, E. W., Monthly Weather Rev., 52, 488-490 (1924).

¹⁶⁵ Holborn, L., Scheel, K., and Henning, F., "Wärmetabellen," Braunschweig, Vieweg & Sohn, 1919.
166 Harrison, L. P., Monthly Weather Rev. 62. 247-248 (1934).

293700,
$$E = -0.0044$$
, $F = -0.0057148$. From (1), (2) is obtained.

$$\frac{dP}{dT} = P \left[-\frac{B + Cx(10^{Dx^2} - 1)}{T^2} + 2C\{10^{Dx^2}(1 + 2Dx^2 \log_e 10) - 1\} - \frac{5}{4}EFy^{1/4} 10^{Fy^{5/4}} \log_e 10 \right] \log_e 10$$
 (2)

Other recent formulas are those proposed by A. Adamson,¹⁵⁷ A. Batschinski,¹⁵⁸ V. Fischer,¹⁵⁹ P. H. Hofbauer,¹⁶⁰ V. Kiréeff,¹⁶¹ and L. B. Smith, F. G. Keyes, and H. T. Gerry.¹⁶²

Certain other formulas constructed for use in interpolating between the directly observed precise values are given in the following tables.

Density and Specific Volume of Saturated Water-vapor.

The pressure, and consequently the density, of the vapor that is in equilibrium with water at a given temperature depends upon the form of the water surface and upon the presence or absence of a foreign gas (p. 559+).

Except where the contrary is stated, the following data refer to pure water-vapor in equilibrium with a flat surface of pure water.

C. H. Meyers ¹⁷⁵ has found that the specific volume (v^*) of a saturated vapor may be calculated, from the vapor-pressure (p) and an approximate value of the specific volume (v') of the liquid, by means of formula (3), p_c being the critical pressure, and A an empirical constant characteristic of the liquid. It fails if $p/p_c > 0.25$. For water, A = 0.651.

$$\log_{10}(1 - pv^*/RT) \cdot (1 - pv'/RT) = A \log_{10}(p/2.718p_c)$$
 (3)

Other interpolation formulas of various types may be found in the sources mentioned in Table 250.

Table 250.—Specific Volume of Saturated Water-vapor

(See also Table 241.)

The published values of the specific volume are here represented each by its defect (δ); *i.e.*, by the amount by which it falls short of v_c^* , defined by the formula $v_o^* = 4.555(273.1 + t)/P_{\rm sat}$, the value of $P_{\rm sat}$ being obtained from Table 242 (except as noted) and 4.555 cm³atm/g.°K being the value accepted by the *International Critical Tables* for the gas-constant (R) of a gas of molecular weight 18.0154 (H_2O). That is, v_c^* is essentially the ideal specific volume for H_2O , the actual specific volume

¹⁶⁷ Washburn, E. W., Int. Crit. Tables, 3, 211-212 (1928).

¹⁸⁸ Prytz, K., Jour. de Physique (2), 3, 353-364 (1893); Math.-fys. Medd. Danske Videnskab. Selskab, 11, No. 2 (1931).

¹⁰⁰ Osborne, N. S., Mech. Eng., 55, 116-117 (1933).

¹⁷⁰ Osborne, Stimson, Fiock and Ginnings, Bur. Stand. Res., 10, 155-158 (RP523) (1933).

¹⁷¹ Osborne, Stimson, and Ginnings, J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937).

¹⁷² Moser, H., Ann. d. Phys. (5), 14, 790-808 (1932).

¹⁷⁸ Koch, W., Forsch. Gebiete Ingenicurw., 5, 257-259 (1934).

¹⁷⁴ Koch, W., Idem, 3, 1-10 (1932).

¹⁷⁵ Meyers, C. H., Bur. Stand. J. Res., 11, 691-701 (RP616) (1933).

Table 250,—(Continued)

being $v^* = v_0^* - \delta$; e.g., at 300 °C the O value for the specific volume is $v^* = 30.786 - 9.142 = 21.644$ cm³/g.

Interpolation is facilitated by making use of the fact that the variation of δ with the temperature is in general small.

With the possible exception of HM, E, and C, each of these sets of data represents the result of an attempt to obtain the most accurate values that can be deduced from all available data. That is, there is a mass of experimental data that is common to them all.

The O values, communicated by Dr. Osborne in May, 1938, are to be preferred.

		Unit	of v_c * ar	ad $\delta = 1$	cm8/g.	Temp.	$= t ^{\circ}C (1$	nt. scale)		
, S	ource"→	0	KSG	JF	HM	Ε δ	IST	M	WT	C	ICT
0 10 20 30 40	206389 106554 57941 33005 19605	103 139 109 78 60	89 144 117 83 62	79 114 101 75 60	289		79 144 117 83 62	68 74 68 61 54	1389 654 281 125 65		
50 60 70 80 90	12093 7719.1 5082.0 3441.2 2390.3	48 41 36 32.6 29.2	48 40.8 35.7 32.0 28.8	48 42 37 33 30	56		48 40.8 35.7 32.0 28.8	46 41 37 32.8 29.4	43 33 32 31 29		
100 110 120 130 140	1699.5 1234.2 913.85 688.76 527.63	26.4 24.0 22.0 20.5 19.0	26.3 24.1 22.20 20.55 19.10	27.4 25.1 23.2 21.5 19.9	27.3		26.3 24.1 22.20 20.55 19.10	26.7 24.3 22.29 20.54 19.04	29		22 26 24.0 21.7 19.5
150 160 170 180 190	410.26 323.43 258.20 208.56 170.28	17.8 16.6 15.57 14.7 13.92	17.80 16.67 15.65 14.76 13.96	18.6 17.3 16.2 15.3 14.4	17.68		17.80 16.67 15.65 14.76 13.96	17.72 16.58 15.56 14.68 13.89			17.9 16.3 15.1 13.7
200 210 220 230 240	140.42 116.86 98.091 82.991 70.732	13.22 12.59 12.02 11.51 11.05	13.24 12.62 12.021 11.508 11.048	13.56 12.84 12.21 11.64 11.14	13.07 11.89		13.24 12.62 12.021 11.508 11.048	13.19 12.56 11.99 11.49 11.03		14.0	
250 260 270 280 290	60 697 52.416 45.532 39.768 34.908	10.64 10.27 9.93 9.63 9.377	10.636 10.267 9.939 9.646 9.386	10.69 10.29 9.93 9.62 9.35	10.572 9.924		10.636 10.267 9.939 9.646 9.386	10.63 10.26 9.92 9.626 9.358		10.41 9.90 9.47 9.12 8.80	
300 310 320 330 340	30.786 27.268 24.246 21.637 19.371	9.142 8.952 8.794 8.669 8.591	9.161 8.968 8.808 8.685 8.607	9.12 8.913 8.751 8.627 8.556	9.192 8.994 8.811 8.640 8.483		9.161 8.968 8.808 8.685 8.607	9.120		8.55 8.36 8.23 8.17 8.19	
350 360 365 366 367	17.391 15.648 14.852 14.698 14.546	8.58 8.71 8.85 8.90 8.96	8.589 8.664	8.554 8.672 8.817		8.598 8.688 8.844	8.685			8.32 8.63	
368 369 370 371 372	14.395 14.246 14.099 13.953 13.808	9.02 9.09 9.17 9.27 9.41	9.006 9.200	9.11 9.33		9.145 9.364	9.192			9.25	
373 374 374.1 374.1 374.2	5 13.500	9.61 10.05 10.4	9.829 10.308			9.892 10.444				9.73	11.08
375 377 380 380.5	13.27d 12.92d 12.27d 11.98d								٠	9.77 9.59 9.44 9.40	

Table 250.—(Continued)

" Sources:

- E
- Callendar, H. L., Proc. Inst. Mech. Eng., 1929, 507-527 (1929).
 Eck., H., Ber. Tätigkeit Phys.-Techn. Reichs. im 1936, p. 32 → Physik. Z., 38, 256 (1937).
 Havlicek, J., and Miškovský, L., Helv. Phys. Acta, 9, 161-207 (1936).
 Int. Crit. Tables, 3, 234 (1928). Compilation by Keyes, F. G., based on work of Knoblauch, O., Linde, R., and Klebe, H., Forsch. Gebiete Ingenieurw., 21, 33-55 (1903), which agrees closely with WT, but with consideration of data by Battelli, A., Ann. chim. et phys. (6), 26, 394-425 (1892); (7), 3, 408-431 (1894), Dieterici, C., Ann. d. Phys. (Wied.), 38, 1-26 (1889), and Perot, A., Ann. chim. et phys. (6), 13, 145-190 (1888).
 International Skeleton Steam Table, 1934, see Table 260.
 Iakol, M., and Fritz, W., Physik. Z., 36, 651-659 (1935). Superseding Jakob, M., Forsch. Geb. Ing., 310, 9-19 (1928), and Jakob, M., and Fritz, W., Z. Ver. deuts. Ing., 73, 629-636 (1929); Techn. Mech. Thermodynam., 1, 173-183, 236-240 (1930); Forsch. Gebiete Ingenicurus., 4, 295-299 (1933).
 Keyes, F. G., Smith, L. B., and Gerry, H. T., Proc. Amer. Acad. Arts Sci., 70, 319-364 (1935). Superseding Smith, L. B., and Keyes, F. G., Mech. Eng., 54, 123-124 (1932).
- IST
- 364 (1935). Superscuing Small, 2. 2., (1932).

 Meyers, C. H., Private communication; values computed by means of formula (3). Osborne, N. S., Private communication, May, 1938. These values supersede all others previously published by Osborne and his associates; in particular they extend to lower temperatures the table by Osborne, Stimson, and Ginnings, J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937) and agree with that table for t>110 °C. Holborn, L., Scheel, K., and Henning, F., "Warmetabellen," 1919.
- ^b From compilation by A. F. O. Germann and S. F. Pickering. 176
- The critical pressure published by Eck (225.5, kg*/cm² = 218.26 atm) was used in the computing of this value.

^d Callendar's value for P_{sat} was used in the computing of this value.

Table 251.—Density of Saturated Water-vapor

To obtain the density of saturated water-vapor, take the reciprocal of the specific volume as given in Table 250, using the particular value of δ that seems most appropriate. The values given in this table were so obtained from the O series, and will serve to indicate the order of magnitude to be expected.

Unit of	$\rho = 1$ g pe	er cm ³ . Temp. = t	°C	
ρ	t	ρ	t	ρ
0.0000048476	80	0.00029338	250	0.019976
0.0000093972	100	0.0005977	300	0.04620
0.0000172912	120	0.0011213	350	0.1135
0.000030370	140	0.001966	360	0.1441
0.000051164	160	0.003259	370	0.203
0.00008302	180	0.005158	374	0.267
0.00013024	200	0.007862	374.15	0.32
	0.0000048476 0.0000093972 0.0000172912 0.000030370 0.000051164 0.00008302	0.0000048476 80 0.0000093972 100 0.000012912 120 0.000030370 140 0.000051164 160 0.00008302 180	0.0000048476 80 0.00029338 0.0000093972 100 0.0005977 0.0000172912 120 0.0011213 0.000030370 140 0.001966 0.000051164 160 0.003259 0.00008302 180 0.005158	0.0000093972 100 0.0005977 300 0.0000172912 120 0.0011213 350 0.000030370 140 0.001966 360 0.000051164 160 0.003259 370 0.00008302 180 0.005158 374

Table 252.—Density of Water-vapor Saturated in the Presence of a Foreign Inert Gas

In the first section of the table, under the title "Ideal case," are given values of ρ'/ρ_0 computed by means of the formula $\log_c(\rho'/\rho_0) = P'/RT$. They are the values that would be obtained were water-vapor an ideal gas, water incompressible and of unit density, and the inert ideal gas insoluble. Departures from these ideal conditions will reduce (see p. 560+) the value of ρ'/ρ_0 , but the effect will be small in all cases covered by this table. In no case will ρ' be less than ρ_0 . These values may serve as a norm against

¹⁷⁶ Germann, A. F. O., and Pickering, S. F., Int. Crit. Tables, 3, 248 (1928).

Table 252.—(Continued)

which to compare the experimental data given in Section II. Marked discrepancies exist.

Pertinent data published by I. R. McHaffie ¹⁷⁷ exhibit strange and unexplained variations with the pressure; they are not included in this table.

For the moisture content of the gas phase in equilibrium with aqueous solutions of NH₃, see Table 233.

 ρ' = mass of water-vapor per liter of actual gas phase; ρ_0 = value of ρ' when no gas except water-vapor is present (its value for any temperature may be found from Table 250); P' = partial pressure of the inert gas; P (or ρ) = total pressure (gas + vapor) corresponding to ρ' .

Unit of P and P'=1 atm = 1.03323 kg*/cm²; of $\rho=1$ kg*/cm²; of ρ' and $\rho_0=1$ mg water-vapor per liter of actual gas phase. Temp. = t °C

1.	Ideal case	(see heading	of table).				
t→ P'	25	37.5	50	$-\frac{100}{\rho'/\rho_0}$	150	200	230
10	1.007	1.007	1.007	1.006	1.005	1.005	1.004
50	1.038	1.036	1.036	1.030	1.026	1.023	1.022
100	1.076	1.073	1.070	1.061	1.053	1.048	1.045
150	1.117	1.112	1.107	1.092	1.081	1.072	1.068
200	1.159	1.152	1.146	1.125	1.109	1.097	1.091
300	1.247	1.236	1.226	1.193	1.168	1.149	1.140
400	1.343	1.327	1.312	1.265	1.231	1.204	1.191
500	1.445	1.424	1.405	1.342	1.296	1.261	1.244
600	1.556	1.528	1.503	1.423	1.365	1.321	1.299
700	1.675	1.640	1.609	1.510	1.438	1.384	1.357
800	1.803	1.760	1.722	1.601	1.515	1.450	1.418
900	1.940	1.889	1.843	1.698	1.595	1.518	1.481

1.801

1.680

1.590

1.547

II. Experimental values.

2.028

2.089

1000

		Ba	rletta, b				
$\begin{matrix} Gas \rightarrow \\ t \rightarrow \\ P \end{matrix}$	H ₂ 50	N ₂ 50	$\frac{50}{\rho'/\rho_0}$	3H ₂ + N ₂ 37.5	25	$Gas \rightarrow t \rightarrow P$	II. 100 \(\rho'/\rho_0\)
100 200 300 400	1.076 1.153 1.206 1.242	1.284 1.551 1.700 1.793	1.130 1.245 1.315 1.368	1.135 1.255 1.345 1.422	1.164 1 337 1.475 1.585	25 50 100 200	1.018 1.04 1.09 1.19
500 600 700	1.276 1.308 1.341	1.876 1.947 1.989	1.425 1.470 1.506	1.481 1.530 1.570	1.662 1.728 1.789	400 600 800	1.40 1.66 1.97
800 900 1000	1.365 1.399 1.431	2.021 2.044 2.053	1.540 1.567 1.582	1.598 1.616 1.643	1.831 1.875 1.909	1000	2.35

1.973

F. Pollitzer and E. Strebel.

Gas→	— Н	o°	A	ire		('O ₂	
$t\rightarrow$	49.9	70.1`	49.9	70.1	$t \rightarrow$	49.9	$t\rightarrow$	70.1
p·		ρ',	/ρ,		Þ	ρ'/ρ_0	p	ρ'/ρ_0
10		1.026	1.011	1.024	40.0	1.747	30.5	1.421
50	0.996	1.041	1.121	1.121	55.5	2.228	39	1.580
100	1.115	1.060	1.258	1.243	59.0	2.597	52.4	1.929
150	1.252	1.154	1.395	1.364	87.0	4.404		

¹⁷⁷ McHaffie, I. R., Phil. Mag. (7), 3, 497-510 (1927).

Table 252.—(Continued)

A. W. Saddington and N. W. K	A.	W.	Saddington	and	N.	W.	Krase."
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Gas→				<i>X</i>	J				_
$P \rightarrow$	100) ——		200			300 -		
ŧ	ρ'	ρ'/ρ_0	ť	ρ'	ρ'/ρ_0	t	ρ'	ρ'/ρ_0	•
50	123.6	1.489	50	146.3	1.763	50	197.5	2.038	
80	381.7	1.302	85	511.6	1.447	75	460.7	1.904	
100	647.4	1.083	150	2960.	1.162	100	912.6	1.527	
150	2710.	1.064	190	7550.	1.180	115	1450.	1.503	
190	5880.	0.919	225	14840.	1.163	145	3240.	1.445	
230	13300.	0.951				165	5320.	1.449	
		0.,,02				230	16400	1.172	

a References:

Bartlett, E. P., J. Am. Chem. Soc., 49, 65-78 (1927); Pollitzer, F., and Strebel, E., Z. physik. Chem., 110, 768-785 (1924); Saddington, A. W., and Krase, N. W., J. Am. Chem. Soc., 56, 353-361 (1934).
W.G. Wiebe, R., and Gaddy, V. L., Idem. 56, 76-79 (1934).

Saturated Liquid

Boiling Point.

The boiling point of a liquid is the temperature at which the vaporpressure of the liquid is equal to the pressure to which the liquid is subjected. It is commonly said to be the temperature of the liquid when it is steadily boiling in such a manner that it is thoroughly intermixed with bubbles of vapor. If the rate of heating is low, if the temperature is measured at a point far from the surface through which the heat is supplied, and if the mixing is ideal and the thermometer is suitably screened from radiation, then the temperature of the boiling liquid approaches closely to the boiling point as defined in the first sentence, and may coincide with it. But under other conditions the temperature of the boiling liquid lies above the true boiling point.¹⁷⁹ The temperature indicated by a suitably screened thermometer immersed in the vapor above the boiling liquid is that at which the vapor at the existing pressure is in equilibrium with the condensed vapor; it is not the same as the boiling point unless the composition of the condensed vapor is the same as that of the boiling liquid. 180

Unless—owing to the presence of nuclei, of submerged solids that are poorly wetted by the liquid and have uneven surfaces penetrated by cavities, or to some other cause—there are in the interior of the liquid free spaces in which the vapor can collect, the liquid can be raised considerably above the boiling point before boiling begins, especially if protected from shocks of all kinds.* F. Donny 182 heated air-free water under a pressure of only a few centimeters of mercury to 135 °C without its boiling. At that tem-

^b Data discussed by J. J. van Laar. ¹⁷⁸

[&]quot;Interpolation by the compiler.

^{*} Since this was written, an important paper by J. Aitken, 181 expressing these same general views, has come to the compiler's attention.

¹⁷⁸ van Laar, J. J., Z. physik. Chem. (A), 145, 207-219 (1929).

¹⁷ Jakob, M., and Fritz, W., Forsch. Gebiete Ingenicurw., 2, 435-447 (1931).

¹⁸⁰ Cf. Schreber, K., Z. techn. Physik, 9, 277-285 (1928).

¹⁸¹ Aitken, J., Trans. Roy. Scottish Soc. Arts, 9, 240-287 (1875).

¹⁸² Donny, F., Ann. de chim. et phys. (3), 16, 167-190 (1846) = Ann. d. Phys. (Pogg.), 67, 562-584 (1846) ← Bruxelles Mém. Couron., 17, (1843-1844).

Table 253.—Boiling Points of Water 156

(Osborne and Meyers)

These values, computed by means of formula (1), are believed to be the best available. Others are compared with these in Table 254.

Each of these values (the temperature at which the saturated vapor exerts the indicated pressure) is the boiling point of water when subjected to the corresponding pressure p exerted by an overlying inert gas when the conditions are such that the boiling has removed from the water all of the previously dissolved gas, and has blanketed the surface of the water with a layer of its own pure vapor. Under other conditions the temperature is somewhat less, lying between this value and that defined by the formula given in the head-matter of Table 245.

Examples: The boiling point when p = 590 mm-Hg is 93.058 °C; when p = 614 mm-Hg it is 94.1319 °C.

Un	it of p	$(= p_1 + p_2)$	$(p_2) = 1$	mm-Hg	= 1.3332	2 milliba	ars. Te	mp. = (t)	$_{1}+t_{2})$ °(C, Int.	scale
$p_2 \rightarrow$		0	1	2	3	4	5	6	7	8	9
P ₁	t_1					100	00 t2 -				
500	88	678	730	782	834	886	938	990	1042	1093	1144
510	89	196	247	298	350	401	452	502	553	604	655
520	89	705	756	806	856	907	957	1007	1057	1107	1157
530	90	206	256	306	355	405	454	503	553	602	651
540	90	700	749	798	846	895	944	992	1041	1089	1138
550	91	186	234	282	330	378	426	474	521	569	617
560	91	664	712	759	806	854	901	948	995	1042	1089
570	92	136	182	229	276	322	369	415	462	508	554
580	92	600	646	692	738	784	830	876	922	967	1013
590	93	58	104	149	195	240	285	330	375	420	465
600	93	510.0	554.8	599.6	644.3	688.9	733.5	778.0	822.4	866.8	911.2
610	93	955.4	999.6	1043.8	1087.9	1131.9	1175.9	1219.8	1263.6	1307.4	1351.1
620	94	394.8	438.4	482.0	525.5	568.9	612.3	655.6	698.9	742.1	785.2
630	94	828.3	871.3	914.3	957.2	1000.1	1042.9	1085.7	1128.4	1171.0	1213.6
640	95	256.3	298.7	341.1	383.4	425.7	468.0	510.2	552.3	594.4	636.5
650	95	678.5	720.4	762.3	804.1	845.9	887.6	929.3	970.9	1012.5	1053.9
660	96	95.4	136.8	178.2	219.5	260.7	301.9	343 1	384.2	425.2	466.2
670	96	507.2	548.0	588.9	629.7	670.4	711.1	751.7	792.3	832.9	873.4
680	96	913.8	954.2	994.6	1034.9	1075.1	1115.3	1155.5	1195.6	1235.6	1275.6
690	97	315 6	355.5	395.4	435.2	474.9	514.6	554.3	593.9	633.5	673.0
700	97	712.5	751.9	791.3	830.7	870.0	909.2	948.4	987.6	1026.7	1065.7
710	98	104.8	143.7	182.7	221.6	260.4	299.2	337.9	376.6	415.3	453.9
720	98	492.5	531.0	569.5	607.9	646.3	684.6	722.9	761.2	799.4	837.6
730	98	875.7	913.8	951.9	989.9	1027.8	1065.7	1103.6	1141.4	1179.2	1217.0
740	99	254.7	292.4	330.0	367.5	405.1	442.6	480.0	517.4	554.8	592.1
750	99	629.4	666.7	703.9	741.0	778.1	815.2	852.3	889.3	926.2	963.1
760	100	. 0	36.8	73.6	110.4	147.1	183.8	220.4	257.0	293.6	330.1
770	100	366.6	403.0	439.4	475.8	512.1	548.4	584.6	620.8	657.0	693.2
780	100	729.3	765.3	801.3	837.3	873.3	909.2	945.0	980.8	1016.6	1052.4
790	101	88.1	123.8	159.4	195.0	230.6	266.1	301.6	337.1	372.5	407.9

Table 254.—Comparison of Values for the Boiling Points of Water

The preferred value is $t_{\rm OM}$ taken from Table 253; each of the other values may be obtained by adding to $t_{\rm OM}$ the corresponding value of $\delta/1000$. For example, the value given by C(V) for the boiling point when p=660 mm-Hg is 96.095-0.013=96.082 °C.

C. S. Cragoe ^{185a} has recommended that, in the absence of more accurate data, the excess, $\Delta t = t' - t$, of the boiling point (t') under pressure p above that (t) under the pressure (A) of one normal atm be computed by

¹⁸⁸ Henrick, F. B., Gilbert, C. S., and Wismer, K. L., J. Phys'l Chem., 28, 1297-1304 (1924).

Table 254.—(Continued)

means of the formula $\phi \Delta t = (273.1 + t') \log_{10}(p/A)$ in which ϕ for a given substance varies but slowly with t', and for any one of the 8 groups, into which he finds all substances for which data are available may be assorted, ϕ varies linearly with the temperature of the normal boiling point. For water at 100 °C, $\phi = 5.79$; it is with this value that the data in column C have been computed.

Other formulas proposed for water may be put in the form t - 100 = $a\pi + b\pi^2 + c\pi^3 + d\pi^4$ where $\pi = (p - 760)/1000$, the pressure being p mm-Hg. Certain of these are included in this table, the values of the coefficients and of the limiting pressures being given at the foot of the appropriate column. Numerous other formulas of the same type have been proposed; some of the more recent ones may be found in papers by L. B. Smith, F. G. Keyes, and H. T. Gerry, 185b J. A. Beattie and B. E. Blaisdell,185c and W. Świętosławski and E. R. Smith.186

U	nit of $p = 1$	mm-Hg = 3	1.33322 mil	libars; of	$\delta = 0.001$	°C; of	$t_{\rm OM}=1~{\rm ^{\circ}C}$	(Int. sca	ile)
Re	ferenceª	Mo	C(V)	ZB	WT	Mu	нн	В	С
<i>†</i> 550 560	^t ом 91.186 91.664		-34 -32		δ- +5 +5		+3		-19 -23
580 600 620	92.600 93.510 94.395		-32 -29 -25 -21		+6 +5 +5		$^{+1}_{+1}$		-10 -10 -9
640 660 680 690 700	95.256 96.095 96.914 97.316 97.712	-4 -5 -3	-17 -13 -10 -9 -6	-5 -6 -5	+4 +4 +2 +2 +3	(+5) +3 +2 +3	$ \begin{array}{c} 0 \\ +1 \\ +3 \\ +1 \\ +2 \end{array} $	+10 +9 +9	-2 -1 -1 0 0
710 720 730 740 750	98.105 98.492 98.876 99.255 99.629	-3 -2 -2 -1 0	-6 -4 -4 -3 -1	-5 -4 -4 -3 -1	+2 +2 +1 +1 +1	+3 +3 +2 +2 +2	0 -1 -1 -1 +1	+7 +6 +4 +3 +2	0 0 0 0
760 770 780 790	100.000 100.367 100.729 101.088	$\begin{pmatrix} 0 \\ 0 \\ 0 \\ (-2) \end{pmatrix}$	0 +1 +3 +4	$0 \\ +1 \\ +2 \\ +3$	0 -1 -1 -1	$ \begin{array}{r} 0 \\ -2 \\ -4 \\ (-8) \end{array} $	$\begin{array}{c} 0 \\ 0 \\ +2 \\ +1 \end{array}$	0 -2 -3 -5	$\begin{array}{c} 0 \\ 0 \\ -1 \\ -1 \end{array}$
o b c d pmin pmax		36.87 -22.00 0 0 690 780	36.970 -19.795 22.83 0 560 820	36.971 -30.263 6.695 0 683 832		36.7 -23.0 0 0 680 780		36.69 -20.45 16.39 -14.3 680 800	9

a References:

Broch, O. J., Trav. et Mém. Bur. Int. Poids et Mes., 1, A43-A48 (1881). Cragoe, C. S. 1858. Volet, C., Trav. et Mém. Bur. Int. Poids et Mes., 18, (1930). (Formula was fitted to P. Chappuis' observations, of which some had not been published before.) Holborn, L., and Henning, F., Ann. d. Phys. (4), 26, 833-883 (1908). 1111

¹⁸⁴ Hoyt, C. S., and Fink, C. L., J. Phys'l Chem., 41, 453-456 (1937).

¹⁸⁵ Rosanoff, M. A., and Dunphy, R. A., J. Am. Chem. Soc., 36, 1411-1418 (1914).

¹⁸⁵a Cragoe, C. S., Inst. Crit. Tables, 3, 246-247 (1928).

Table 254.—(Continued)

Mo	Moser, H., Idem (5), 14, 790-808 (1932).
Mu	Formula given by Mueller, E. F., in Int. Crit. Tables, 1, 53 (1926) and recom-
	mended by the International Bureau of Weights and Measures in 1927 in connec-
	mended by the litternational bureau of weights and measures in 1927 in connec-
	tion with the International Temperature Scale. See: C. R. des Septième Conf.
	Gén. Poids et Mes., 1927, 56-58, 94-99 (1928). Burgess, G. K., Bur. Stand. J.
	Res., 1, 635-640 (RP22) (1928).
WT	D. 1 6 m 11 04ff 6 (431/2 11 11 11
VV I	Derived from Table 245, from "Wärmetabellen."
7 D	7 1 A T T A C 1 400 4000 1000 (1000) 4 T
ZB	Zmaczynski, A., and Bonhoure, A., Compt. rend., 189, 1069-1070 (1929) ← Jour.
	do Phase (7) 1 295-201 (1020)

de Phys. (7), 1, 285-291 (1930).

perature the vapor-pressure of water is over 3 atm. The initiation of boiling under such conditions results in an explosion of the entire volume of water. F. B. Kenrick, C. S. Gilbert, and K. L. Wismer 183 report that they have heated water in a capillary tube and at atmospheric pressure to 270 °C.

Likewise, in the absence of condensation nuclei, a vapor can be cooled, without condensation, below the temperature corresponding to equilibrium between it and its liquid (see p. 633).

Effect of a Solute.—The boiling point of a solution exceeds that of the pure solvent by an amount Δ , and we may write $\Delta = nE$, where n is the number of effective gram-molecules contributed by the solute per kilogram of the solvent. E is characteristic of the solvent but independent of the nature of the solute if that is not significantly volatile at the boiling point; it varies with the concentration, and its limiting value (E_0) as n approaches zero is frequently called the ebullioscopic constant of the solvent. If the solute is neither associated nor dissociated, and does not affect the molecular aggregation of the solvent, then $n = 1000m/FW \equiv N$, where m is the mass of the solute dissolved in the mass W of the solvent, and F is the molecular weight of the solute. If each molecule of the solute is dissociated into two parts, and if the solvent is not affected, then n = 2N; and similarly in other cases. The value of E_0 is $RM_wT^2/1000L$, where R is the gasconstant (=8.315 joules/g-mole. $^{\circ}$ K), M_w = molecular weight of the vapor of the solvent, $T \circ K =$ absolute temperature of the boiling point of the solution, and L = latent heat of vaporization of the pure solvent at $T \circ K$. (R and L must refer to the same unit of mass.) For very dilute aqueous solutions, $T=373.1\,^{\circ}\text{K}$, $L=2256.6\,\text{j/g}$, and E_0 for water is $8.315M_w$ × $(373.1)^2/1000(2256.6M_w) = 0.513$ °C per (effective g-mole of solute per kg of water). Experimental data agree with this value as well as one should expect.

(C. S. Hoyt and C. L. Fink 184 state that the best values now in use for the ebullioscopic constants are those published by M. A. Rosanoff and R. A. Dunphy. 185 Those authors, using $v^* = 1.651$ liters/gram for the specific volume of water-vapor saturated at 100 °C, obtained $E_0 = 0.518$ for water. But the best value of v^* now available is 1.673 (see Table 250); had they used that, retaining the same values as before for the other con-

¹⁸⁵⁶ Smith, L. B., Keyes, F. G., and Gerry, H. T., Proc. Amer. Acad. Arts Sci., 69, 137-168 (1934).

¹⁸⁵c Beattie, J. A., and Blaisdell, B. E., Idem, 71, 361-374 (1937).

¹⁸⁶ Swietosławski, W., and Smith, E. R., J. Res. Nat. Bur. Stand., 20, 549-553 (RP1088) (1938).

stants, they would have found $E_0 = 0.511$. Hoyt and Fink, using $R = 1.976 \times 4.185 = 8.270$ joules/g-mole. K, obtained $E_0 = 0.510$ for water. Had they used the value (R = 8.315) adopted by the *International Critical Tables*, and used in this compilation, retaining the other constants unchanged, they would have found $E_0 = 0.513$.)

Density and Specific Volume of Saturated Water.

The best published extended series of values of the specific volume of saturated water is that contained in the International Skeleton Steam

Table 255.—Specific Volume of Saturated Water

(See also Table 241.)

The preferred values (v^*) are given in full, adjacent to the t-column; for other values merely the excess (δ) of each above the preferred value is given, the unit of δ being that of the last place in the tabulated value of v^* . For example, the ICT value for 120 °C is 1.0603-0.0011=1.0592; and Eck's value at the critical point is 3.1-0.034=3.066. The difference between the significance of the decimal points (in the δ values and in the v^* values) should be remembered.

The pressure is always that of the saturated vapor.

Unit of $v^* = 1$ cm³/g; of $\delta = 1$ unit in the last tabulated place in v^* ; of p = 1 kg*/cm². Temp. = t °C (Int. scale)

				_					
I.	Temperat	ure give	en.						
Source	a→ IST	SK		IST = SK	ICT		OSG	Eck	IST
t	v^*	δ	ŧ	v^*	δ	t	v^*	δ-	
0	1.00021	+2	100	1.0435	-1	330	1.562		-0.1
10	1.00035	+21	110	1.0515	-5	340	1.640		+0.8
20	1.00184	+20	120	1.0603	-11	350	1.741	+4	+5.8
30	1.00442	+11	130	1.0697	-17	355	1.808	+2	
40	1.00789	+2	140	1.0798	-22	360	1.894	0	+12.6
50	1.0121	0	150	1.0906	-25	365	2.016	-2	
60	1.0171	-1	160	1.1021	-24	366	2.048		
70	1.0228	$-2 \\ -1$	170	1.1144	-20	367	2.083		
80	1.0290		180	1.1275	-13	368	2.124		
90	1.0359	-1	190	1.1415	-5	369	2.170		
100	1.0435	-1	200	1.1565	+4	370	2.225	-14	+6
		_	210	1.1726	+13	371	2.293		+4
			220	1.1900	+19	372	2.38	-3.3	+0.1
			230	1.2087	+13	373	2.51		-0.8
			240	1.2291	$+1_{9}$	374	2.80	-15.0	-1
			250	1.2512	$+0_{8}$	374.15	3.1	-0.34^{b}	
			260	1.2755	-1_{5}				
			270	1.3023	—5₃				
			280	1.3321					
			290	1.3655					
			300	1.4036					
			310	1.4475					
			320	1.4992					
II.	Pressure	given.	Sª						
		50	_	100	150		200	22	25
	v.	1.286		1.451	1.649		1.990	3.0	

Table 255.—(Continued)

" Sources:

Eck, H., Ber. Tatigkeit Phys.-Techn. Reichs. in 1936, p. $32 \rightarrow Physik. Z.$, 38, 256 (1937). Eck

(1937).

Int. Crit. Tables, 3, 234 (1928). Compiled by Keyes, F. G., and based on the data of Hirn, G. A., Ann. Chim. et Phys. (4), 10, 32-92 (1867), Ramsay, W., and Young, S., Phil. Trans. (A), 183, 107-130 (1892), and Waterston, J. J., Phil. Mag. (4), 26, 116-134 (1863).

International Skeleton Steam Table, 1934 (see Table 260). The values below 100 °C have been derived from the Chappuis and the Thiesen values (see text); those for t = 100 to 360 °C are the same as the SK values.

Osborne, N. S., Stimson, H. F., and Ginnings, D. C., J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937); confirmed by Osborne in May, 1938.

Schlegel, E., Z. techn. Phys., 14, 105-107 (1933).

Smith, L. B., and Keyes, F. G. 188 This paper supersedes that of Keyes and Smith, 180 and of Smith and Keyes, Mech. Eng., 55, 114-116 (1933). ICT

IST

OSG

^b This is for Eck's value at the critical point, which he places at 374.23 °C. In their compilation ¹⁸⁹ A. F. O. Germann and S. F. Pickering give 2.5 for the critical volume; and by extrapolation Keyes and Smith ¹⁹⁰ found 3.086 cm³/g.

Table of 1934,¹⁸⁷ given in Table 260. For temperatures below 100 °C they have been derived from the determinations at one atm by Chappuis and by Thiesen (see Table 93) by applying a suitable correction for the isothermal compression from saturation to a pressure of one atm, and by changing the unit of volume from 1 ml to 1 cm³. From 100 to 360 °C they are identical with those of the extended table published by L. B. Smith and F. G. Keyes, 188 except for a rounding off to a smaller number of digits. The Smith and Keyes table was computed by means of an empirical formula (4) fitted to their observations above 100 °C and taking into consideration known values for lower temperatures.

$$v^* = \left[v_o + a(t_o - t)^{1/3} + b(t_o - t) + c(t_o - t)^4\right] / \left[1 + d(t_o - t)^{1/3} + e(t_o - t)\right] \text{ cm}^3/\text{g}$$
(4)

where $v_c = 3.197500$, $t_c = 374.11$ °C, a = -0.3151548, b = -0.001203374, $c = +7.48908/10^{13}, d = +0.1342489, e = -0.003946263.$

For temperatures above 330 °C the slightly different values given by Osborne, Stimson, and Ginnings (1937, OSG of Table 255) are to be preferred.

For the density take the reciprocal of the specific volume.

89. THERMAL ENERGIES OF SATURATED WATER AND SATURATED STEAM

In this section are given values of the enthalpy (heat content) and of the entropy of both saturated water and saturated steam, and the specific heat of steam continuously saturated. The quantity frequently called the specific heat at saturation really refers to the unsaturated state, being merely the limit approached by the specific heat of the unsaturated phase as the temperature approaches that of saturation. Its values are not given here,

¹⁶⁷ See Mech. Eng., 57, 710-713 (1935).

¹⁸⁸ Smith, L. B., and Keyes, F. G., Proc. Amer. Acad. Arts Sci., 69, 285-314 (1934) → Mech. Eng., 56, 92-94 (1934).

¹⁸⁹ Germann, A. F. O., and Pickering, S. F., Int. Crit. Tables, 3, 248 (1928).

¹⁸⁰ Keyes, F. G., and Smith, L. B., Mech. Eng., 53, 132-135 (1931).

but will be found in the corresponding sections treating of the unsaturated phases.

For a recent review of the experimental data, see E. F. Fiock.¹⁹¹

Table 256.—Enthalpy of Saturated Water and of Saturated Steam

The OSG values to to be preferred. They are given directly, the others by the amounts (ΔII) by which they exceed the corresponding OSG values. Example: The value given by KSG for the saturated vapor at 0 °C is 2500.00 + 1.86 = 2501.86.

II is the excess of the enthalpy (E + pv) above its value for saturated water at 0 °C. Subscripts l and v indicate that the value refers to the liquid and to the vapor, respectively.

Unit of H and $\Delta H=1$ Int. joule/g. Temp. = t °C (Int. scale) Values adopted for International Skeleton Steam Tables, 1934, are given in Table 260.

	000	~	— Liquio	TC.	C-1-		OCC		Vapo		77	
Ref.ª→	H_{i}	0	K	ΤS ΔΗ ι	Sch	С	OSG Hv	0	KSG	HM H _v	K	C
0	0.000	0	0.00	0.00	0.0	0.00	2500.00	0	+1.86	+0.3		•
10	42.028	ŏ	0.00	0	+0.1	0.00	2518.50	ŏ	+1.61	1 0.0		
10 20	83.833	Ō	-0.03	- 0.16	+0.1		2536.81	Ö	+1.43			
30	125.675	Õ			0.0		2554.97	Ō	+1.30			
40	167.454	ŏ	-0.09	-0.01	-0.4		2572.99	Ŏ	+1.18			
50	209.247	ŏ	-0.16	0.01	-0.2		2590.83	ŏ	+1.03	-0.9		
60	251.072	ŏ	-0.20	+0.09	-0.3		2608.44	ŏ	+0.83	0.5		
70	292.943	ŏ	0.20	. 0.05	-0.2		2625.77	ŏ	+0.62			
80	334.877	ŏ	-0.29	0.00	-0.3		2642.75	ŏ	+0.58			
90	376.893	ő	0142	0.00	-0.4		2658.31	ŏ	+0.18			
100	418.76	0.25	-0.16	- 0.16	0.0		2675.42	-0.14	-0.17	-1.9		
110	460,99	0.26	0.10	0.10	0.0		2690.85	+0.05	-0.20	1.5		
20	503.34	0.31	-0.2	1 0.24	0.0		2705.56	+0.22	-0.23			
130	545.94	0.29	0.2	1 0.24	0.6		2719.62	+0.28	-0.32			
40	588.73	0.29	-0.2	+ 0.45	0.3		2732 05	1 0.29	-0.45			
50	631.76	0.30	- 0.1	1 0.75	0.1		2732.95 2745.49	+0.30	-0.65	-0.7		
60	675.07	0.30	+0.1	+ 0.76	-0.1		2757.18	+0.31	-0.94	0.7		
70	718.69	0.31	10.1	1 0.70	-0.1		2767.87	+0.31	-1.28			
.80	762.68	0.32	0.0	+ 0.76	-0.6		2777.29	+0.31	-1.48			
90	807.07	0.32	0.0	7 0.70	-0.6		2785.35	+0.32	-1.54			
90	851.91	0.32	+0.8	√ 0.14	-1.4	+0.5	2791.99	+0.32	-1.54 -1.53	100		**
10	897.27	0.33	TU.8	7 0.14	-1.4	0.8	2797.11	+0.33	-1.33	+0.9		10 12
220	943.21	0.33	+0.8	-0.32	-2.7	1.3	2800.62	+0.32	-1.40	+2.3		14
230	000 00	0.32	TU.0	0.32	-3.6	1.6	2802.38	+0.32	-1.25	T2.3		14
	989.80		+0.2	-0.77					-1.25			16
40 50	1037.13 1085.30	0.32	+0.2	-0.77	-6.1	2.0	2802.25	+0.33		100		17
	1134.44	0.32	+0.1	0.00	-8 -10	2.4 2.7	2800.07 2795.61	+0.33	-1.07 -1.03	+3.3		19 21
260		0.32	0.0	-0.88		2.7		+0.33		100		21
70	1184.69	0.33	-0.5	1075	-15	3.5 4.2	2788.61	+0.32	-1.04	+2.2		23
80	1236.25	0.32	-1.4	1 2.75	-18	4.2	2778.72	+0.33	-1.09			25
90	1289.35	0.33	4.0		-23	4.8	2765.52	+0.32	-1.19			27
00	1344.29	0.32	-1.8	+ 4.41	- 29	5.5 6.5	2748.38	+0.32	-1.33	-1.1	-4.0	29
10	1401.50	0.32	-2.1		-36	6.5	2726.42	+0.32	-1.36	-1.3		31
20	1461.56	0.32	-1.9		-36	7.5	2698.81	+0.32	-1.49	-0.9		32
30	1525.36	0.32	-2.1		-38	8.6	2664.65	+0.32	-2.14	+1.0		33.
40	1594.23	0.31	- 2.2		-34	9.8	2621.23 2563.42	$+0.32 \\ +0.31$	-2.79	+6.8		33
50	1670.56	0.32	-4.6		-32	11.3	2563.42	+0.31	-0.846		-5.3	34
60	1760.86	0.32	-1.9		-30	18.1	2480.28	+0.33	+10.998		-2.2	38
65	1816.88	0.32					2420.12	+0.33				
66	1829.72	0.32					2405.42	+0.31				
167	1843.38	0.32					2389.42	+0.32				
68	1858.08	0.32					2371.84	+0.31				
68	1873.99	0.32					2352.23 2329.99	+0.33				
70	1891.69	0.32	-0.9		-20	28.8	2329.99	+0.32			-0.5	67
71	1911.88	0.31					2303.98	+0.33				
72	1936.11	0.32	-1.3				2272.15	+0.32			-1.6	
73	1968.21	0.33					2229.19	+0.32				
174	2031.07	0.33			+60	-13.50		+0.34				174.
374.15	2083.27	0.32					2083.27	0.32				

¹⁹¹ Fiock, E. F., Bur. Stand. J. Res., 5, 481-505 (RP210) (1930) → Mech. Eng., 52, 231-242 (FSP-52-30) (1930).

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Osborne, N. S., Private communication, 1938. (These values take precedence over the corresponding OSG ones.)

Osborne, N. S., Stimson, H. F., and Ginnings, D. C. Values for t < 100 have been privately communicated by Osborne, 1938, and are subject to slight revision as the work progresses; the others are from a longer table in J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937). These supersede similar values published by Osborne, Stimson, and Fiock, Bur. Stand. J. Res., 5, 411-480 (Rl'209) (1930) \rightarrow Mech. Eng., 52, 191-220 (FSP-52-28) (1930) and by Osborne, Stimson, and Ginnings, Mech. Eng., 56, 94-95 (1934); Idem, 57, 162-163 (1935). OSG

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b These values were obtained by KSG by extrapolation.

° Callendar, holding that the true critical temperature is 380.5 °C, extends his observations to higher temperatures: 275 °C, $H_v = 2296.9$, $H_1 = 2051.5$; 377 °C, $H_v = 2242.0$, $H_1 = 2055.8$; 380 °C, $H_v = 2077.5$; $H_1 = 2029.3$; and 380.5 °C, $H_2 = H_1 = 1968$.

Table 257.—Specific Heat of Saturated Water-vapor 192

 c_s = amount of heat per g that must be added in order to increase the temperature by 1 °C while the condition of saturation is maintained by suitably adjusting the volume.

Table 258.—Entropy of Saturated Water and of Saturated Steam

(Values adopted for the International Steam Tables, 1934, are given in Table 260.)

The OSG values are to be preferred. They are given directly, the others by the amounts (ΔS) by which they exceed the corresponding OSG values. Example: For the saturated vapor at 0 °C M gives the equivalent of $S_v = 9.132 - 0.0037 = 9.1283$.

The values under C and M illustrate the magnitude of the errors in the tables current ten years ago.

S is the excess of the entropy above its value for saturated water at $0 \,^{\circ}$ C. The subscripts l and v indicate that the corresponding values refer to the liquid and to the vapor, respectively.

¹⁹² Leduc, A., Int. Crit. Tables, 5, 83 (1929).

Table 258.—(Continued)

Unit of S and of $\Delta S = 1$ Int. joule/g. °C. Temp. = t °C (Int. scale)

	_	- Liquid			- Vapor -	_
Ref.a→	OSG Sı	K C 104ΔS	М	OSG	C 104	AS- M
0	0	0	0	9.132	10	-37
10	0.1511		+1	8.884		69
20	0.2962	+2	+3	8.656		-84
30 40	0.4363	-1	4 5	8.446°		-82 -78
	0.5719	1		8.253		
50 60	0.7032 0.8305	0	11 11	8.074 7.909		59 34
70	0.9543	U	17	7.756		$-34 \\ -15$
80	1.0746	0	15	7.613		+12
90	1.1918		7	7.480		+40
100	1.3059	-24^{b}	9	7.3536		+85
110	1.4174		12	7.2371		+107
120 130	1.5265 1.6332	+6	14 19	7.1278 7.0248		+128
140	1.7378	+7	20	6.9276		$+149 \\ +162$
150	1.8404		24	6.8355		+166
160	1.9412	- 1 7	25	6.7480		+170
170	2.0403		27	6.6643		+161
180	2.1380	+ 11	25	6.5836		+156
190	2.2342		26	6.5055		+141
200	2.3293	+15 +23 +25	18	6.4295	+245	+118
210 220	2.4233 2.5164	+15 + 36	11 +2	6.3554 6.2827	270 297	+84 +45
230	2.6088	+38	-14	6.2112	326	-10
240	2.7007	+1 +48	-33	6.1404	351	-77
250	2.7922	+46	-60	6.0699	382	-159
260	2.8835	-6 + 62	-93	5.9992	415	-252
270	2.9749	+78 -18 +88	-137 -193	5.9279	441	-351
280 290	3.0668 3.1594	-18 + 88 + 100	-260	5.8553 5.7807	473 498	-463 -575
300	3.2534	-29 + 123	-334	5.7032	524	-696
310	3.3495	-36 + 121	-399	5.6214	542	-799
320	3.4482	-39 + 139	-448	5.5341	639	-869
330	3.5504	-36 + 151	-498	5.4392	581	-912
340	3.6583	-35 + 190	-539	5.3332	586	-882
350	3.7763	-68 + 245	-605	5.2091	591	-797
360 365	3.9150 4.0002	+306 +375	-568	5.051 <i>2</i> 4.9455	634 728	-658
366	4.0002	+3/3		4.9203	120	
367	4.0401			4.8930		
368	4.0620			4.8634		
369	4.0858			4.8305		
370	4.1121	+471	-123	4.7936	1062	-506
371	4.1422			4.7509		
372	4.1787			4.6695		
373	4.2282	-247	+902	4.6321 4.5079	2663	869
374	4.3308	-247	+ 902	4.30/9	2003	009

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C K

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Table 258.—(Continued)

more extended table by Oshorne, Stimson, and Ginnings, J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937) which takes precedence over the former.

^b Koch's published value (0.3114 Int. steam cal/g) for the saturated liquid at 100 °C seems to involve a misprint; if it were 0.3119, then $10^4\Delta S_1$ would be -3joules/g.°C, which is not very discordant with the adjacent values.

° From fundamental constants and spectroscopic data, A. R. Gordon and Colin Barnes 108 computed the excess of the entropy of saturated steam at 30 °C above that at absolute zero to be 54.39 cal/g-mole(°C) = 12.642 j/g(°C).

STEAM-TABLES AND DIAGRAMS

Although many steam-tables and diagrams, each giving several types of steam-engineering data pertaining to one or more of the systems—water, steam, water and steam—have been published, there was in 1929 no generally accepted consistent set of data from which complete steam-tables could be constructed. Tables based on different sources differed disconcertingly, although it is probable that any one of several was good enough for most technical purposes, provided its data were not mixed with those from another source. Any mixing had to be done with circumspection.

As a result of this condition, an international steam-table conference was held in London in 1929, another in Berlin in 1931, and a third in this country in 1934. A skeleton steam-table was adopted at the first, and was revised and somewhat extended at the second; and a more extended one was adopted at the third. The last, which supersedes all others, is here given as Table 260. A review of the better work on steam prior to the London conference has been published by H. N. Davis and J. H. Keenan. 194

In Table 259 are listed some of the more recent steam tables and diagrams and of the reports of extended work pertaining thereto. Extracts from much of the latter may be found distributed through the several sections devoted to the properties concerned.

The calorimetric measurements in this field have been recently reviewed by E. F. Fiock 195; and many pertinent data based on measurements made at the Physikalisch-Technischen Reichsanstalt are given by L. Holborn, K. Scheel, and F. Henning in their "Wärmetabellen" (1919).

Numerous equations of state have been proposed for water-vapor, but none has been generally accepted as entirely satisfactory. Some will be found in the references given elsewhere in this section, and some are discussed in the following papers, listed in chronological order: Linde, R., Forsch. Gebiete Ingenieurw., 21, 57-92 (1905). Eichelberg, Idem, 220, 1-31 (1920), Tumlirz, O., Sitz. Akad. Wiss. Wien (Abt. 2a), 130, 93-133 (1921), Strauven, M., Rev. univers. des mines (6), 16, 289-301, 363-376 (1923), Callendar, H. L., World Power, 1, 274-280, 325-328 (1924), Nesselmann, K., Z. physik. Chem., 108, 309-340 (1924), Hausen, H., Forsch, Gebiete Ingenieurw., 2, 319-326 (1931).

¹⁰⁸ Gordon, A. R., and Barnes, Colin, J. Phys'l Chem., 36, 1143-1151 (1932).

¹⁹⁴ Davis, H. N., and Keenan, J. H., Proc. World Eng. Cong., Tokyo, 1929, 4, 239-264 (1931). ¹⁸⁸ Fjock, E. F., Bur. Stand. J. Res., 5, 481-505 (RP210) (1930) = Mech. Eng., 52, 231-242 (FSP-52-30) (1930).

Table 259.—A List of Some Recent Steam Tables and Diagrams and of Reports of Extended Work Pertaining Thereto

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Table 260.—International Skeleton Steam-Tables, 1934

The following values were adopted by the Third International Steam-Tables Conference held in the United States of America a in 1934, 196 and involve the following units and conversion factors: Inch, pound, atmosphere, and bar, all as already defined in Table 1; the international steamtable calorie, designated as "IT-cal" defined by the relation: 1000 IT-cal = 1/860 international kilowatthour; 1 international watt = 1.0003 (absolute) watt; 1 British thermal unit = 251.996 IT-cal; temperatures are expressed in terms of the international (centigrade) temperature scale, the zero being regarded as equivalent to 273.16 °K. These lead to the following equivalents: $1 \text{ kg*/cm}^2 = 14.2233 \text{ lb*/in}^2 = 0.967841 \text{ atm}$; $1 \text{ cm}^3 = 3.53146_7$ 10^{-5} ft³; 1 m³/kg = 16.0185 ft³/lb; 1 IT-cal = 4.18605 Int. j = 21.447 10^{-3} lb*·ft³/in² = 41.3255 atm·cm³ = 4.1873 joules.

The enthalpy or total heat is defined as "the heat content in excess of that contained by the liquid at zero degree centigrade and saturation pressure."197

 $P = \text{pressure}, \ v^* = \text{specific volume}, \ H = \text{enthalpy (total heat)}; \ \text{sub-}$ scripts l and s indicate that the data refer to the liquid and to the vapor, respectively; $\tau =$ tolerance, expressed in units of the last place of the tabulated value: Tol. is the actual tolerance, for the entire column of values above it, unless another tolerance is indicated.

Observed values $t_{crit} = 374.11$ °C (M.I.T.), 374.2 ± 0.1 °C (P.T.R.).

Unit of $P = 1 \text{ kg*/cm}^2$; of $v^* = 1 \text{ cm}^3/\text{g}$; of H = 1 IT-cal/g; of $\tau = 1 \text{ unit in last place}$.

I. Saturated phases: Liquid (1), vapor (s).

b	P	τ	vi^*	τ	v * *	τ	$^{\prime}H\iota$	τ	H_s	τ
0	0.006228	6	1.00021	5	206310	210	0	0	597.3	7
10	0.012513	10	1.00035	10	106410	110	10.04	1	601.6	7
20	0.023829	20	1.00184	10	57824	58	20.03	2	605.9	6
3 0	0.043254	30	1.00442	10	32922	33	30.00	2	610.2	5
40	0.075204	38	1.00789	10	19543	19	39.98	2	614.5	5

¹⁹⁰ Mech. Eng., 57, 710-713 (1935).

¹⁹⁷ London Conf.; Mech. Eng., 52, 120-122 (1930).

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Table	200	-1 601		eu,

				abic 20	o. (Commin					
	P	τ	v_l*	τ	v.*	τ	H_{l}	τ	H_{\bullet}	7
50 60 70 80 90	0.12578 0.20312 0.31775 0.48292 0.71491	6 10 16 24 36	1.0121 1.0171 1.0228 1.0290 1.0359	2 2 2 2 2	12045 7678.3 5046.3 3409.2 2361.5	12 77 50 34 24	49.95 59.94 69.93 79.95 89.98	3 3 4 5	618.9 623.1 627.3 631.4 635.3	5 5 5 5 5 5
100 110 120 130 140	1.03323 1.4609 2.0245 2.7544 3.6848	0 10 13 16 21	1.0435 1.0515 1.0603 1.0697 1.0798	2 4 4 4 4	1673.2 1210.1 891.65 668.21 508.53	17 12 89 67 51	100.04 110.12 120.25 130.42 140.64	5 6 7 7	639.1 642.7 646.2 649.6 652.7	5 5 5 5 6
150 160 170 180 190	4.8535 6.3023 8.0764 10.225 12.800	32 42 53 7 8	1.0906 1.1021 1.1144 1.1275 1.1415	4 4 4 4	392.46 306.76 242.55 193.80 156.32	39 31 24 19 16	150.92 161.26 171.68 182.18 192.78	8 8 9 9	655.7 658.5 661.0 663.3 665.2	7 8 8 9 9
200 210 220 230 240	15.857 19.456 23.659 28.531 34.140	8 9 10 12	1.1565 1.1726 1.1900 1.2087 1.2291	4 4 4 4 4	127.18 104.24 86.070 71.483 59.684	13 10 86 71 60	203.49 214.32 225.29 236.41 247.72	10 11 11 12 12	666,8 668,0 669,0 669,4 669,4	9 9 9 9
250 260 270 280 290	40.560 47.866 56.137 65.457 75.917	13 15 17 20 22	1.2512 1.2755 1.3023 1.3321 1.3655	4 4 4 4 5	50 061 42.149 35.593 30.122 25.522	50 42 36 30 30	259.23 270.97 282.98 295.30 307.99	13 18 19 20 20	668.9 667.8 666.0 663.6 660.4	9 9 9 9
300 310 320 330 340	87.611 100.64 115.12 131.18 148.96	24 3 3 4 4	1.4036 1.4475 1.4992 1.5619 1.6408	55555	21.625 18.300 15.438 12.952 10.764	35 35 35 35 35	320.98 334.63 349.00 364.23 380.69	30 40 50 60 70	656 1 650.8 644.2 636.0 625.6	10 12 14 16 18
350 360 370 371 372	168.63 190.42 214.68 217.26 219.88	4 5 5 10 11	1.7468 1.9066 2.231 2.297 2.381	6 40 21 26 34	8.802 6.963 4.997 4.761 4 498	35 40 100 100 110	398.9 420.8 452.3 457.2 462 9	8 15 15 22	611.9 592.9 559.3 553.8 547.1	20 20 30 35 40
373 374	222.53 225.22	11 11	2.502 2.79	53 15	4.182 3.648	120 120	471.0 488.0	35 50	538.9 523.3	45 50

II. Compressed water.

$\stackrel{t\rightarrow}{P}$	0	50	100	150	200	250	300	350
1	1.00016	1.01210						
5	0.9999	1.0119	1.0432	1.0906				
10	0.9997	1.0117	1.0431	1.0902				
25	0.9989	1.0110	1 0422	1.0893	1 1556			
50	0.9977	1.0099	1.0409	1.0877	1.1532	1.2495		
75	0.9965	1.0088	1.0397	1.0861	1.1508	1.2452		
100	0.9952	1.0077	1.0385	1 0845	1.1485	1 2410	1.3979	
125	0.9940	1.0067	1.0372	1 0829	1.1462	1.2369	1.3877	
150	0.9929	1.0056	1.0360	1.0814	1.1439	1.2330	1.3782	
200	0.9905	1.0035	1.0337	1.0784	1.1395	1 2255	1.3612	1.671
250	0.9882	1.0015	1.0314	1.0755	1 1353	1.2184	1.3462	1.604
300	0.9859	0.9995	1.0291	1.0726	1.1312	1.2117	1.3327	1.557
350	0.9837	0.9975	1.0269	1.0698	1.1272	1.2054	1.3207	1.521
400	0.9814	0.9956	1.0247	1.0670	1.1234	1.1994	1.3097	
Tol.→	0.0002	0.0002	0.0002	0.0002	0.0003	0.0004	0.0007	0.002

Table 260.—(Continued)

$\stackrel{t\rightarrow}{P}$	0	50	0 100		150 ————————————————————————————————————	200	250	30	00 3	350
1 5 10 25 50 75 100 125 150 200 250 300 Tol.	0.023 0.120 0.240 0.599 1.20 1.79 2.98 3.57 4.74 5.90 7.05 → 0.01°	49.5 50.0 50.1 50.4 50.9 51.4 51.9 52.4 52.9 53.9 54.9 0.0	100.1 15 100.2 15 100.4 100.9 16 101.3 101.7 16 102.2 16 102.6 103.5 104.4 105.3	20 66 60 44 82 57 65	150.92 151.00 151.21 151.58 151.95 152.32 152.32 153.06 153.82 154.57 155.33 0.08	203.6 203.8 204.1 204.3 204.6 204.8 205.2 205.8 206.2 0.1	259.2 259.2 259.2 259.3 259.3 259.4 259.5 259.7 0.1 4	320 319 318 312 312	9.9 9.3 3.4 39 7.6 38 7.0 38	93.1 17.6 14.0 0.8
11			(i.e., dila	ted)	steam.					
$\stackrel{t\rightarrow}{P}$	v.* 10	00	v.* 150	T	7', * 200	7	v. * 250 -	T	v.* 300	T
1 5 10 25 50 75	1730	1	1975	2	2215 433.8 210.4	2 4 2	2454 484.1 237.6 89.0	2 5 2 1	2691 533.2 263.3 101.1 46.41 27.48	3 5 3 1 7 5
$P \xrightarrow{t \to P}$	7',* 350 -		7's* 400 -	_	~** 45	0 —	~ 500		v ₃* 55	0 —
1 5 10 25 50 75 100 125 150 200 250 300	2928 581.6 288.2 112.1 53.12 33.22 23.03 16.66 11.98	3 6 3 1 8 7 5 3 2	3164 629.6 312.7 122.6 59.05 37.78 27.05 20.53 16.10 10.31 6.366 3.02	3 6 3 1 9 8 5 4 3 1 3 1	3400 677.4 337.0 132.7 64.60 41.83 30.41 23.52 18.90 13.05 9.46 6.98	8 6 5 4 3 2 2	3636 725.0 361.1 142.7 69.92 45.62 33.45 26.14 21.25 15.11 11.39 8.90	4 7 4 1 10 9 7 5 4 3 2 2	3872 772.5 385.1 152.6 75.10 49.25 36.32 28.55 23.36 16.87 12.96	4 4 4 2 12 10 7 6 5 3 3 2
t→ P 1 5 10 25 50 75	100 - 11s 639.2	5	663.2	τ 5	686.5 681.9 675.1	6 10 10	710.1 706.7 702.1 687.8	6 10 11 11	734.0 731.5 728.0 718.0 698.4 672.6	12 12 12 12 12 12
t→ 1 5 10 25 50 75 100 125 150	758.0 756.1 753.5 746.3 732.9 717.6 699.5 676.7 646.8	12 12 12 12 12 12 12 12 12 12 12 15	782.4 782.4 780.8 778.9 773.3 763.1 752.1 740.0 726.9 712.1	12 12 12 12 12 12 12 13 13 13	H, 45 807.2 805.9 804.5 800.0 791.6 783.2 774.5 765.2 755.3	12 12 12 12 12 12 12 12 13 13 13	832.3 831.3 830.1 826.5 819.9 813.1 806.0 799.1 791.8	12 12 12 12 12 12 12 12 13 13 13	H _s 550 857.8 856.9 855.9 852.6 847.3 841.8 836.1 830.3 824.4	20 20 20 20 20 20 20 20 20 20 20 20

Table 260.—(Continued)

$t\rightarrow$	350		H. 40	0	H. 450)	H ₀ 500		550	
$\stackrel{t\rightarrow}{P}$	H_{\bullet}	τ`	H_{\bullet}	τ	H_{\bullet}	7	H_{\bullet}	τ .	H _e	τ
200			676.5	20	733.4	20	<i>77</i> 6.0	20	812.0	25
250			622.5	25	707.5	25	758.8	25	798.9	30
300			524.5	30	677.5	25	739.7	25		

^a The first conference was held in London, Engineering (London), 128, 751-752 (1929), Mech. Eng., 52, 120-122 (1930), Z. Vev. Deuts. Ing., 73, 1856-1858 (1929); the second, in Berlin, Z. Ver Deuts. Ing., 75, 187-188 (1931), Mech. Eng., 53, 287-290 (1931), Engineering (London), 131, 296-297, 393 (1931).

^b For t=0 and P=1, Tol. = 0.00005. ^c For t=0 and P=1, 5, 10, and 25, Tol. = 0.005. ^d For P=250 and t=200 and 250, Tol. = 0.2; for P=300 and t=200 and 250, Tol. = 0.3.

91. FUGACITY AND ACTIVITY OF WATER

The value of the fugacity of water, as that term is used by G. N. Lewis and M. Randall, 198 is $f = P_s e^{-(g-w)}$ where P_s is the pressure of the vapor when saturated at the temperature $(T \circ K)$ considered, and g and w are the values of two definite isothermal integrals, one (g) determined solely by the vapor (gas), and the other (w) solely by the liquid (water); RTg =

$$\int_{P_0}^{P_s} (\delta/P) dP \text{ and } RTvv = \int_{P_s}^{P'} v_w * dP, \text{ where } \delta = RT - Pv *, v \text{ being}$$

the specific volume of the vapor at T $^{\circ}$ K and under the pressure P, P_0 is any such low pressure that it is reasonable to assume that δ is negligible if $P \equiv P_0$, v_w^* is the specific volume of water at the same temperature and under the pressure P, P' is the pressure at which the fugacity is desired, and R is the gas-constant. Obviously, the units of v^* , v_w^* , and P must be so chosen that the unit of each of the products Pv^* and v_w^*dP , and also of δ , shall be the same as that of the RT with which it is associated.

The expression for w may conveniently be broken into two parts, one involving integration from P_s to A, the other from A to P', where A denotes the pressure of 1 atm. These integrals we shall denote by w' and w'', respectively;

$$RTw' = \int_{P_*}^{A} v_w dP$$
, $RTw'' = \int_{A}^{P'} v_w * dP$, $w = w' + w''$.

If P' = A, w'' = 0 and the fugacity at 1 atm is $f_1 = P_s e^{-(y \cdot w')}$; if $P' = P_s$, w' = -w'' and the fugacity at saturation is $f_s = P_s e^{-g}$. Also $f = P_s e^{-(g-w'-w'')} = f_1 e^{w''}$; Lewis and Randall call the quantity $e^{w''}$ the activity and denote it by a; thus getting the relation $f = af_1$.

Values for f and for a of water at 25, 37.5, and 50 °C, and at various pressures between 1 and 1000 atm have been published by M. Randall and B. Sosnick. 199 Those values are quoted by the former in his compilation, 200

Lewis, G. N., and Randall, M., See their "Thermodynamics and the Free Energy of Chemical Substances," 1923. Also, Tunnell, G., J. Phys'l Chem., 35, 2885-2913 (1931).
 Randall, M., and Sosnick, B., J. Am. Chem. Soc., 50, 967-980 (975) (1928).

²⁰⁰ Randall, M., Int. Crit. Tables, 7, 232 (1930).

but are affected by computational errors and an assumption. These affect two of the 5 significant digits given in the values of f, and should be carefully considered.

Their procedure was to compute f_1 and a, and to derive $f(=af_1)$ from them. Any error or uncertainty in f_1 will recur, in full force, in each other value of f for that temperature; on that, will be superposed any error in the corresponding value of a.

As there seem to be no suitable experimental data for the density of dilated water-vapor at these temperatures, the value of g can be obtained only by the use of an assumed equation of state. They used Berthelot's

equation, taking
$$\delta = \frac{9}{128} \left(\frac{RPT_c}{P_c} \right) \left\{ 6 \left(\frac{T_c}{T} \right)^2 - 1 \right\}$$
, and took for the critical

constants $P_c = 217.5$ atm, $T_c = 647.1$ °K, essentially those given in the *International Critical Tables* and in this compilation. As extrapolation always introduces an uncertainty, it is imperative that we consider carefully the probable suitability of this equation for our present purposes.

Probably the best test is to examine how well this equation will reproduce, at such temperatures, the known density of the saturated vapor. If the reproduction is close, it is probable that the equation is fairly satisfactory; in the contrary case, and especially if the difference between the two values increases rapidly as the temperature is reduced, the equation must be condemned as unsatisfactory, and the results of computations based upon it must be regarded as of doubtful value. The quantities of interest in such a comparison are given in Table 261, where also are given those computed by means of a new equation of state, derived from observations at and above 100 °C, and taking into consideration the specific heat as well as the density. That equation ²⁰¹ yields

$$\delta = \left(\frac{54.8}{v^*} - \frac{42.6}{v^{*2}}\right)T - \frac{81.8(10)^3}{v^*} + \frac{38.7(10)^6}{v^*T} + \frac{32.5(10)^{28}}{v^{*2}T^9} \text{cm}^3 \text{atm/g}$$

It will be noticed (Table 261) that at the higher temperatures (50 to $200\,^{\circ}$ C) the values derived from Naumann's equation do not differ greatly from those given by the observations; while those from Berthelot's are only about half as great as the experimental values at $50\,^{\circ}$ C, and 3/4 as great at $200\,^{\circ}$ C. At lower temperatures, both sets depart markedly from the experimental values, the Berthelot values at $25\,^{\circ}$ C being only 1/5 as great as the experimental. The Naumann value always lies between the other two. Whence, we may conclude that neither of these equations of state is suitable for use in computing g for temperatures below $50\,^{\circ}$ C, and that Berthelot's is quite unsuitable even at $50\,^{\circ}$ C.

The conservative course is to restrict oneself to the statement that g is probably near, and does not exceed, the value given in the column Obs. For such small quantities we may for our present purposes take $e^{-g} = 1 - g$. Then the expression for the fugacity of water at 1 atm becomes $f_1 = 1 - g$.

²⁰¹ Naumann, F., Z. Physik Chem. (A), 159, 135-144 (1932).

 $P_{se}^{-(g \ w')} = P_s(1-g)c^{w'}$, where 1000g does not exceed 3.0, 2.4, and 3.4, respectively, at 25, 37.5, and 50 °C. At those same temperatures, $c^{w'} = 1 + \tau v'$, where $1000\tau v' = 0.72$, 0.67, and 0.60, respectively. Whence at 50 °C, for example, $f_1 = P_s(1.00060 - g) \ge P_s(1.00060 - 0.0034) = 0.9972P_s$. Putting into the expression for f_1 the values of P_s (0.031258, 0.063637, and 0.12172 atm) corresponding, respectively, to 25, 37.5, and 50 °C, we find that f_1 is near, and not less than 0.03119, 0.06353, and 0.12138 atm, respectively. It seems certain that f_1 is not greater than the value derived from Berthelot's equation; vis., 0.03126, 0.06361, and 0.12158. Hence, one may say that at 25, 37.5, and 50 °C, the values of f_1 lie within the ranges 0.03119 to 0.03126, 0.06353 to 0.06361, and 0.12138 to 0.12158 atm, respectively, and probably much nearer the lower values.

Table 261.—A Comparison of Some Bases for an Estimate of the Fugacity of Saturated Water

(See text for discussion, symbols, and formulas.)

The fugacity at saturation is
$$f_s = P_s e^{-\eta}$$
, $RTg = \int_{P_0}^{P_s} (\delta/P) dP$, $\delta \equiv$

 $RT - Pv^*$, $\Delta \equiv \delta/P$, v^* is the specific volume of the vapor; subscript s indicates that the quantity refers to the saturated vapor. B indicates that the value is based on Berthelot's equation of state, N on Naumann's, Obs. on observed data.

Unit of vs*	and $\Delta_s = 1$	cm ⁸ /g; of δ_{κ}	= 1 cm ³ atm/g	g is	dimensionless.	Temp. = t °C
		Α.			8	104

		Δ,				δ			-104a -	
<i>‡</i>	v_s *	Óbs.	N	B`	Obs.	N	в`	Obs. a	Ň	B
0	205000	1400	61	31	8.9	0.37	0.19	71	3	1
10	105900	522	54	28	6.3	0.65	0.34	51	5	3
20	57660	20_{6}	46	27	4.8	1.1	0.62	36	8	5
25	43310	13 ₀	47	26	4.1	1.4	8.0	30	11	6
30	32880	9_1	46	25	3.8	1.0	1.0	28	14	8
37.5	22178	54	40	24	3.4	2.5	1.5	24	18	11
50	12050	41	38	22	50	3.1	2.7	34	32	18
100	1671	28	26	16	28.	26.	16.	168	149	96
140	507.7	19.8	20.0	13.1	70.6	71.4	46.7	376	363	248
200	127.1	13.4	14.3	9.8	206.	<i>22</i> 0.	150.	951	924	730

[&]quot;Here it is assumed that $\Delta = \Delta$, at all values of P less than P. Observations above 100 °C indicate that Δ is almost independent of P.

It will be noticed that the values here given as derived from Berthelot's equation differ from the published values to which reference has been made. That is because in obtaining those values the wrong sign was inadvertently given to g. That error affects the entire table.

The values for the activity (a) which are given by Randall and Sosnick are said to have been derived from the values of v_w^* found by P. W. Bridgman.²⁰² But those data, being given only at intervals of 500 kg*/cm² in the pressure, are not entirely satisfactory for use in the defermination

²⁰² Bridgman, P. W., Proc. Am. Acad. Arts Sci., 48, 307-362 (1913).

of the integral w''. Using a linear interpolation over each of the two intervals (1 to 500, and 500-1000 kg*/cm²), which will give a value somewhat too great, the graph of v^* vs. P being convex toward the axes, the present compiler obtained from those data values of a that, at the higher pressures, were significantly less (18 and 24 units in the last place at 1000 atm and at 25 and 50 °C, respectively) than those given in the table cited. This indicates an error in those tabulated values. He did not attempt to check the values at 37.5 °C.

Table 262.—Activity and Fugacity of Water at 50 °C

At 50 °C the fugacity of water is $f = P_s(1.00060 - g)a$, where $P_s =$ pressure of the vapor at saturation, a = activity of water at the pressure considered, $1.00060 = e^{w}$, and g is determined by the extent to which the behavior of the vapor departs from that of an ideal gas. It is probable that for water-vapor at 50 °C, g is close to, and does not exceed, 0.0034. Taking $P_s = 0.12172$ atm, f = 0.12179(1 - g)a atm.

	L'nit	of P and of f - 1	atm; a is dim	ensionless	
P	а	f/(1-g)	P	а	f/(1-g)
1	1	0.12179	600	1.5018	0.18290
100	1.0703	0.13035	700	1.6058	0.19557
200	1.1459	0.13956	800	1.7165	0.20905
300	1.2266	0.14939	900	1.8345	0.22342
400	1.3126	0.15986	1000	1.9601	0.23872
500	1.4042	0.17102			

Consequently, no values from the table cited are tabulated in this compilation, but values of a for water at 50 °C have been derived from Amagat's data (Table 95), which are available in steps of 50 atm, and are given, with the corresponding value of f, in Table 262. They differ by a maximum of 3 units in the last place from those obtained by the compiler from Bridgman's data, linear interpolation being used.

92. Pressure-temperature Associations for Equilibrium Between an Ice and Another Phase

In this section are assembled all those pressure-temperature associations that are characteristic of each of the several types of ice when in such equilibrium as can exist between it and pure water-vapor, pure water, and each of the other types of ice. For convenience, a few values of the specific volume and density of water-vapor in equilibrium with ice are also given. For the density of water and ice, see Sections 32 and 67.

Triple Points.

The triple points of a substance are the temperatures at which three phases of it can coexist in equilibrium. To each such temperature corresponds a unique pressure.

Table 263.—Triple Points of the Water-substance

(Adapted from the compilation by P. W. Bridgman ICT.²⁰³)

The symbols in the first column indicate the three phases that are in equilibrium at the temperature and pressure indicated in the next two columns; G = vapor, L = liquid (water), I, II, III, V, VI, and VII indicate, respectively, six types of ice.

	Unit of P	= 1 atm.	Temp. = t °C	
	Phases		t	P
G	I	$\Gamma_{\mathfrak{b}}$	+0.0099	0.00603
I	L	III	-22.0	2047
I	II	III	-34.7	2100
II	III	V	-24.3	3397
III	L	v	-17.0	3417
V	L	VI	+0.16	6175
VI	L	VII	+81.6	21700 204

^{*}Based on Bridgman, P. W., Proc. Am. Acad. Arts Sci., 47, 439-558 (1912); Z. anorg. Chem., 77, 377-455 (1913); Z. physik. Chem., 89, 252-253 (1915); with a consideration of Tammann, G., "Kristallisieren and Schmelzen," 1903; Z. physik. Chem., 84, 257-292 (1913); 88, 57-62 (1914); "Aggregatzustände," 1922.

*See p. 604 (P=4.58 mm-Hg = 0.00603 atm), also head text of Table 245 (P=4.5867 mm-Hg = 0.0060351 atm).

Ice and Water-vapor

The only type of ice that can exist in equilibrium with water-vapor is the usual one, ice-I. At each temperature below 0 °C there is a unique pressure of the vapor at which equilibrium exists between icc-I and the That pressure is called the vapor-pressure of ice, or the saturation pressure of water-vapor in contact with ice-I. At other pressures one of these phases grows at the expense of the other without the intervention of the liquid phase—ice sublimes. But certain observations by H. T. Barnes and W. S. Vipond 205 suggest that the vapor from ice is for a brief interval in a state somewhat different from its permanent one (see p. 000).

Vapor-pressure of Ice-I.—By integrating the Clausius-Clapeyron equation $(dp/dt = L/(v^* - V^*)T)$ and determining the several constants from certain selected data, mentioned later, E. W. Washburn 208 derived formula (1).

$$\log_{10} p = -\frac{2445.5646}{T} + 8.2312 \log_{10} T - 1677.006(10^{-6}) T + 120514(10^{-10}) T^2 - 6.757169$$
 (1)

Here p mm-Hg is the vapor-pressure of ice-I at t °C, and T = 273.1 + t. The coefficient of the first term of the right-hand member is determined by the value of the latent heat of ice at 0 °C, and was derived from C. Dieterici's value 208a for the latent heat of vaporization of water, and the value

- 200 Bridgman, P. W., Int. Crit. Tables, 4, 11 (1928).
- 204 Bridgman, P. W., Chem'l Phys., 5, 964-966 (1937).
- 205 Barnes, H. T., and Vipond, W. S., Phys. Rev., 28, 453 (1909).
- 200 Washburn, E. W., Monthly Weather Rev., 52, 488-490 (1924).
- 208a Dieterici, C., Ann. d. Physik (Wied.), 37, 494-508 (1889).
- ²⁰⁷ Dickinson, H. C., and Osborne, N. S., Bull. Bur. Stand., 12, 49-81 (S248) (1915).

found by H. C. Dickinson and N. S. Osborne 207 for the latent heat of fusion of ice, both at 0 °C; that of the second term was derived from Dickinson and Osborne's 207 value for the specific heat of ice-I, and upon an estimated value (0.457 cal₂₀/g·°C) for the specific heat of water vapor, both at 0 °C; those of the third and fourth, from two experimental values of p: That found by S. Weber 208 at -100 °C, and that by K. Scheel and

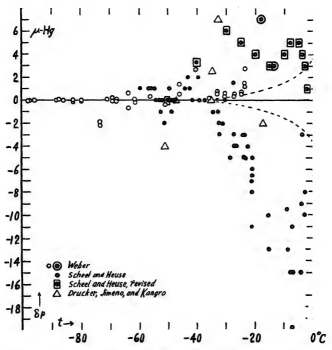


FIGURE 12. Vapor Pressure of Ice-I: Deviations of Observations from the Values Defined by Formula (1).

[Adapted from E. W. Washburn, Monthly Weather Rev., 52, 488-490 (1924).]

δρ = ρ_{eale} - ρ_{obs}; unit of δρ = 1 μ·Hg = 0.001 mm·Hg. The dotted curves indicate the change produced in δρ by an error of 0.01 °C in the temperature.

Weber = Weber, S., Com. Phys. Lab. Leiden, No. 150, 3-52 (1915).

Scheel and Heuse = Scheel, K., and Heuse, W., Ann. d. Physik (4), 29, 723-737 (1909).

Scheel and Heuse, revised = the preceding as revised by Holborn, L., Scheel, K., and Henning, F., "Wärmetabellen," 1919.

Drucker, Jiméno, and Kangro = Drucker, C., Jiméno, E., and Kangro, W., Z. physik. chem., 90, 513-552 (1915).

W. Heuse 209 at -50 °C; and the constant term was so determined as to make p take the well-established value 4.579 mm-Hg at 0 °C.

Washburn was of the opinion that the experimental values in the range -50 °C to near 0 °C are unreliable and should be rejected, the errors arising perhaps from uncertainties in the temperatures. See Fig. 12.

C. F. Marvin 210 has discussed the experimental determinations made prior to 1909.

Table 264.—Vapor-pressure of Ice-I

(Adapted from a table given by E. W. Washburn,²¹¹ the values being those defined by formula (1). See text and Fig. 12. Values below -99 °C have been computed by the compiler.)

 p_s = pressure of pure water-vapor that is in equilibrium with pure ice-I at t °C. If the vapor is mixed with atmospheric air, total pressure being P, then the pressure of the vapor when equilibrium exists will be $p = p_s + \Delta p$, where $\Delta p = V^*(P - p)p_s/RT$, V^* being the volume of unit mass of ice at T °K, and R being the gas-constant. For all practical purposes this is equivalent to $100 \Delta p/p_s = 24/T$ when P = 1 atm.†

	Unit of $p_s = 1$ mm-Hg. Temp. = $t ^{\circ}$ C = $(t_1 + t_2) ^{\circ}$ C										
$t_2 \rightarrow t_1$	-0.0	-0.1	-0.2	-0.3	-0.4	-0.5	-0.6	- 0.7	- 08	-0.9	
-0	4.579	4.542	4.504	4.467	4.431	4.395	4.359	4.323	4.287	4.252	
-1 -2 -3	4.217	4.182	4.147	4.113	4.079	4.045	4.012	3.979	3.946	3.913	
-2	3.880	3.848	3.816	3.785	3.753	3.722	3.691	3.660	3.630	3.599	
-3	3.568	3.539	3.509	3.480	3.451	3.422	3.393	3.364	3 336	3.308	
-4	3.280	3.252	3.225	3.198	3.171	3.144	3.117	3.091	3.065	3.039	
-5	3.013	2.987	2.962	2.937	2.912	2.887	2.862	2.838	2.813	2.790	
	2.765	2.742	2.718	2.695	2,672	2 649	2.626	2.603	2.581	2.559	
-6 -7	2.537	2.515	2.493	2.472	2.450	2,429	2.408	2.387	2.367	2.346	
$-8 \\ -9$	2.326	2.306	2.285	2.266	2.246	2.226	2.207	2.187	2.168	2.149	
-9	2.131	2.112	2.093	2.075	2.057	2.039	2.021	2.003	1.985	1.968	
-10	1.950	1.934	1.916	1.899	1.883	1.866	1 849	1.833	1.817	1.800	
-11	1.785	1.769	1 753	1.737	1.722	1.707	1.691	1.676	1.661	1.646	
-12	1.632	1.617	1.602	1.588	1.574	1.559	1.546	1.532	1.518	1.504	
-13	1.490	1.477	1.464	1.450	1.437	1.424	1.411	1.399	1.386	1.373	
-14	1.361	1.348	1.336	1.324	1.312	1.300	1.288	1.276	1.264	1.253	
-15	1.241	1.230	1.219	1.208	1.196	1.186	1.175	1.164	1.153	1.142	
-16	1.132	1.121	1.111	1.101	1.091	1.080	1.070	1.060	1.051	1.041	
-17	1.031	1.021	1.012	1.002	0.993	0.984	0.975	0.966	0.956	0.947	
-18	0.939	0.930	0.921	0.912	0.904	0.895	0.887	0.879	0.870	0.862	
-19	0.854	0.846	0.838	0.830	0.822	0.814	0.806	0.799	0.791	0.783	

	Unit of p	e = 0.001 mm-I	Ig. Temp. $= t$	$^{\circ}$ C = $(t_1 + t_2)$ $^{\circ}$ C	
$t_2 \rightarrow t_1$	-0	-1	2	-3	-4
-20	776.	705.	640.°	580.	526.
-30	285.9	257.5	231.8	208.4	187.3
-40	96.6	86.2	76.8	68.4	60.9
-50	29.5 ₅	26.1	23.0	20.3	17.8
-60	$8.0_{\rm s}$	$7.0_{\rm a}$	6.1.	5.34	4.64
-70	1.94	1.67	1.43	1.23	1.05
80	0.40	0.34	0.29	0.24	0.20
90	0.07_{0}	0.05_{8}	0.04_{8}	0.04_{o}	0.03_{n}
-100	$0.009_{\rm p}$	0.008_{1}	0.006₅	0.005_{3}	$0.004_{\rm R}$
-110	0.0010	$0.0008_{\rm s}$	0.0006_{0}	0.0005₅	0.0004

[†]This coefficient (24) differs from that (20) given by Washburn 211; the latter is incorrect.

²⁰⁸ Weber, S., Comm. Phys. Lab. Leiden, No. 150, 3-52 (1915).

²⁰⁰ Scheel, K., and Heuse, W., Ann. d. Physik (4), 29, 723-737 (1909).

²¹⁰ Marvin, C. F., Monthly Weather Rev., 37, 3-9 (1909).

³¹¹ Washburn, E. W., Int. Crit. Tables, 3, 210-211 (1928)

Monthly Weather Rev., 52, 488-490 (1924).

Table	264	-(Continued)
Lauic	204	- Committee

$t_2 \rightarrow$	- 5	-6	-7	-8	-9
-20	476.	430.	389.	351.	317.
-30	168.1	150.7	135.1	120.9	108.1
-40	54.1	48.1	42.6	37.8	33.4
-50	15.7°	13.8	12.1	10.6	9.2_{8}
-60	4.0 _թ	$3.4v^a$	3.0₂	2.6_{1}	2.25
-70	0.90	0.77	0.66	0.56	0.47
-80	0.17	0.14	0.12	0.10	0.084
-90	0.02_{7}	0.02_{2}	$0.01_{\rm s}$	0.015	0.01_{2}
-100	0.0034	$0.002_{\rm s}$	0.002_{2}	0.001	0.001
-110	0.0003₄	0.0002_{6}	0.0002_{o}	0.0001	0.0001_{2}

^a C. Dei ²¹⁸ has obtained experimentally the values $p_s = 0.58 \pm 0.023$ mm-Hg at -22.3, 0.0166 ± 0.0008 at -55, and 0.0037 ± 0.0002 at -66 °C, which essentially agree with those (0.62, 0.0157, 0.0035) given by this table.

^b S. Dushman's estimate ²¹² of p_s at -110 °C is 0.75×10^{-6} mm-Hg, somewhat smaller than the 1.0×10^{-6} required by Washburn's equation.

Density and Specific Volume of Vapor Saturated with Respect to Ice. At such low vapor-pressures, the vapor may for most practical purposes be regarded as ideal. Its ideal specific volume at t $^{\circ}$ C may be computed

Table 265.—Ideal Specific Volume and Density of Vapor Saturated with Respect to Ice

Computed by means of the formula $v^* = 3461.8(273.1 + t)/p_s$ cm³/g, where v^* = specific volume, p_s mm-Hg = pressure of vapor in equilibrium with ice-I at t °C (see Table 264); ρ = density of the vapor.

Unit of $v^* = 1$ m ³ /g; of $\rho = 1$ mg/m ³												
ŧ	v*	ρ	t	10 ⁻³ v*	100ρ							
-0	0.2065	4843	-60	0.1427	701							
$-0 \\ -5$	0.3195	3129	-65	0.290	344							
-10	0.5026	1990	-70	0.612	166							
-15	0.8036	1244	<i>−7</i> 5	1.34	75							
-20	1.308	765	-80	3.06	33							
-25	2.168	461	-85	7.3	14							
-30	3.670	272.5	-90	18.0	5.6							
-35	6.345	157.6	-95	47	2.1							
-40	11.22	89.1	-100	130	0.77							
45	20.36	49.1	-105	385	0.26							
-50	37.85	26.42	-110	1330	0.075							
-55	72.3	13.8	-115	3900	0.025							

by means of the relation $v^* = 4.555(760)(273.1 + t)/p_s = 3461.8 \times$ $(273.1 + t)/p_s$ cm³/g, the unit of p_s being 1 mm-Hg, and the values of p₈ (the vapor-pressure of ice) being taken from Table 264. But the actual specific volume is somewhat smaller than the ideal. A tentative determination by M. Knudsen 214 indicated that the effective molecular weight

²¹⁸ Dushman, S., Int. Crit. Tables, 1, 92 (1926).

²¹⁸ Dei, C., Atti acc. Lincei (6), 12, 119-124 (1930).

²¹⁴ Knudsen, M., Ann. d. Physik (4), 44, 525-536 (1914).

of water-vapor saturated at -75 °C is 21.1, which corresponds to a specific volume about 15 per cent less than the ideal.

Ice and Water.

Melting-point of Ice

By the melting-point of ice is meant the temperature at which ice is in equilibrum with the adjacent water under the existing pressure. It varies with the pressure and with the purity of the water.

Except as the contrary is indicated, the following data refer to pure ice in contact with pure water, the system being subjected to a uniform hydro-

Table 266.—Absolute Temperature of the Ice-point

The value that was used for computational purposes in the *International* Critical Tables, and that is so used in this compilation except where another is indicated, is this: Ice-point = $0 \, ^{\circ}\text{C} = T_0 \, ^{\circ}\text{K} = 273.1 \, ^{\circ}\text{K}$. But it is quite certain that the correct value of T_0 is somewhat greater than that, probably about 273.16 °K. For a recent discussion of the subject, see KT.^a

	Unit of $T_0 = 1$ °K										
Year	T_{0}	Ref	Year	T_0	Refa						
1907	273.174 ^b	Buckingham	1930	273.16	НО						
1915	273.09	Henning	1931	273.116 d	Keyes						
1919	273.09°	HSH	1931	273.16	KK						
1921	273.20	HH	1932	273.16°	KS						
1922	273.135 ± 0.005	Keyes	1933	273.215	Jacyna						
1923	273.13 ± 0.01	ST	1934	273.16	KSG						
1925	273.15	Roebuck	1935	273.22	Jacyna						
1929	273.14 ± 0.01	Schames	1936	273.144	KT						
1929	273.16	HO	1936	273.16	Roebuck						
1929	272.79 ^a	Roebuck									

a References:

References:
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Henning, F., "Die Grundlage...der Temperaturmessung," p. 80, Vieweg, 1915.
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Henning, F., and Heuse, W., Z. Physik, 5, 285-314 (1921).
Heuse, W., and Otto, J., Ann. d. Physik (5), 2, 1012-1030 (1929); 4, 778-780 (1930).
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Keyes, F. G., Smith, L. B., and Gerry, H. T., Mech. Lng., 56, 87-92 (1934). Also, Smith, L. B., Keyes, F. G., and Gerry, H. T., Proc. Am. Acad. Arts Sci., 69, 137-168 (1934).
Kessom, W. H., and Tuyn, W., Comm. K. Onnes Lab. Leiden, Suppl. 78, 1-85 (1936) ← Trav. ct Mem. Bur. Int. Poids ct Mes., 20, (1936).
Roebuck, J. R., Proc. Am. Acad. Arts Sci., 60, 537-596 (1925); 64, 287-334 (1929); Phys. Rev. (2), 50, 370-375 (1936).
Schames, L., Z. Physik, \$7, 804-807 (1929).
T. Smith, L. B., and Taylor, R. S., J. Am. Chem. Soc., 45, 2124-2128 (1923).
From a consideration of his own and preceding determinations, Buckingham HH HŠH KK KS KSG

From a consideration of his own and preceding determinations, Buckingham concluded that 273.13 "is probably not far from the true value."

This value of T₀ was derived from the value (0.0036618) given in the "Wärme-

tabellen" (p. 20) for the coefficient of expansion of an ideal gas.

d The computation by which Roebuck obtained the value 272.79 has been criticized by Keyes, who by a recomputation from the same data obtained the value 273.116.

In the paper here referred to, Keyes and Smith suggested that T_0 "may not be far from 273.16."

static pressure. For the effect of differential stresses, see Sections 64 (pp. 430-431) and 97 (p. 646). Five types of ice (I, III, V, VI, and VII) must be considered.

Ice-I: Normal Melting-point and Triple-point.—By definition, the melting-point of ice-I (ordinary ice) in contact with water saturated with air

Table 267.—Melting-points of the Ices under Hydrostatic Pressure

Adapted from the compilation by P. W. Bridgman (ICT)^{a, 221} with the addition of dP/dt from his paper, ²²² (1912) and of such other data as are so indicated. The data for ice-V and ice-VI extend into the supercooled regions. For phase diagram, see Section 93.

Unit of P = 1 atm = 1.01325 bars = 1.0332 kg*/cm². Melting-point = t °C dP/dt P ŧ dP/dt Ice-I - Ice-VI -+0.0099^a -15.0+96.40.00603 4640 5110 +103.00.0 -134.3-10.0590 -5.0-112.05620 -5.0+110.21090 -10.0-95.46160 0.0 +116.2-15.0-82.2+0.16°1540 6175 1910 -20.0-71.86780 +5.0+121.8 -22.0° 10.0 +128.52047 7390 653 -5.53^{t} 8040 15.0 +135.58710 20.00 831 -7.46° +143.8-9.75'1064 10250 30.0 +162.0-10.42'11990 40.0 +182.71104 -12.74^{\prime} 50.0 +208.71313 13970 -15.66^{t} 16150 60.0 +235.11546 18530 70.0 21090 80.0 Ice-VIIh Ice-III -+1802047 -22.0°21700 81.64 108.4 -20.095.3 121.6 2430 +23823200 25200 2820 -18.5+310110.3 134.8 -17.0^{d} 144.5 3417 +43027100 124.1 137.1 152.7 29000 - Ice-V -149.5 31000 161.2 3040 -200+119161.1 32900 170.6 -17.0^{4} 3417 34800 172.1 180.5 3680 -15.0+13536800 182.5 190.8 4360 +152-10.038700 192.3 202.0 5270 -5.0+1746160 0.0 +2016175 +0.16

^a Based on P. W. Bridgman ²²² with a consideration of G. Tammann. ²²³

^b Triple point (vapor, water, ice-I); see Table 263. The increase (dT) produced in the melting point of ice by a small increase (dP) in the hydrostatic pressure can be computed by Clapeyron's equation: $dT/dP = T(\Delta v^*/L)$, where T = absolute temperature of the melting-point, $\Delta v^* =$ excess of the specific volume of water (the phase obtained by increasing the temperature) over that of ice, and L = latent heat of

²¹⁵ Thomson, J., Trans. Roy. Soc. Edinburgh, 16, 575-580 (1849); Thomson, W. (Lord Kelvin), Phil. Mag. (3), 37, 123-127 (1850).

²¹⁶ Moser, H., Ann. d. Physik (5), 1, 341-360 (1929).

²¹⁷ Michels, A., and Coeterier, F., Proc. Akad. Wet. Amsterdam, 30, 1017-1020 (1927).

²¹⁸ Goossens, B. J., Arch. néerl., 20, 447-454 (1886).

Table 267.—(Continued)

fusion; all at the temperature T. For ice at 0 °C, T=273.1 °K, $\Delta v^*=0.0906$ cm³/g, L=333.6 joules/g; whence, dT/dP=-0.00742 °K/bar = -0.00752 °K/atm. H. Moser ²²⁴ observed -0.00748 °K/atm. There is a further depression of 0.0024 °K/atm due to the solubility of air in water.

- Triple point: water, ice-I, ice-III.
 Triple point: water, ice-III, ice-V.
 Triple point: water, ice-V, ice-VI.
 Observations by G. Tammann.²²⁵
- ° L. H. Adams 220 found that water and ice-VI are in equilibrium at 25 °C if P=9630 bars =9504 atm.
 - AP. W. Bridgman 227; units changed by the compiler.

'Triple point: water, ice-VI, ice-VII.

at a pressure of one normal atmosphere, but otherwise pure, the entire system being subjected to a uniform hydrostatic pressure of one normal atmosphere, is 0 °C. This is often called the normal melting-point of ice; also, the ice-point.

The effect of the dissolved air is to lower the melting-point by 0.0024 °C (Table 268), and the direct effect of the pressure (1 atm) is to lower it by 0.0075 °C,²¹⁵ see Table 267. Hence, if it were possible to reduce the pressure to zero, then the melting-point of ice-I in contact with pure, airfree water would be +0.0099 °C. Actually, the pressure cannot be reduced below that of the vapor-pressure of water at the temperature of equilibrium, which must equal the vapor-pressure of ice at the same temperature. That pressure is equivalent to 4.58 mm-Hg, which corresponds to a negligible (0.00004₅ °C) lowering of the melting-point. Consequently, the temperature at which the system (water, vapor, ice-I) is in equilibrium is +0.0099 °C; this is called its triple-point.

H. Moser ²¹⁶ has reported that the triple-point is experimentally reproducible to within 0.0005 °C, and has recommended that it, rather than the ice-point, be used for fixing the 0 °C point of thermometric instruments. See also A. Michels and F. Coeterier ²¹⁷ and B. J. Goossens. ²¹⁸ But J. L. Thomas ²¹⁹ of this Bureau has found that the ice-point can be "readily reproduced to a few ten thousandths of a degree centigrade," and W. P. White ²²⁰ has described procedures by which it can be reproduced and held constant to within 0.0001 °C for a day or more.

Effect of a Solute.—When an aqueous solution, other than a cutectic mixture, begins to freeze, the crystals that form are pure ice (see pp. 401+

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<sup>216</sup> Thomas, J. L., Bur, Stand. J. Res., 12, 323-327 (RP658) (1934).
<sup>220</sup> White, W. P., J. Am. Chem. Soc., 56, 20-24 (1934).
<sup>221</sup> Bridgman, P. W., Int. Crit. Tables, 4, 11 (1928).
<sup>222</sup> Bridgman, P. W., Proc. Am. Acad. Arts Sci., 47, 439-558 (1912) → Z. anorg. Chem., 77, 377-455 (1913); Z. physik. Chem., 89, 252-253 (1915).
<sup>223</sup> Tammann, G., "Aggregatrustande." 1922; "Krystallisieren und Schmelzen," 1903; Z. physik. Chem., 84, 257-292 (1913); 88, 57-62 (1914).
<sup>224</sup> Moset, JI., Ann. d. Physik (5), 1, 341-360 (1929).
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²²⁴ Mosei, H., Ann. d. Physik (5), 1, 341-360 (1929). ²²⁵ Tammann, G., Z. physik. Chem., 72, 609-631 (1910).

²²⁰ Adams, L. H., J. Am. Chem. Soc., 53, 3769-3813 (1931).

²²⁷ Bridgman, P. W., J. Chem'l Phys., 5, 964-966 (1937).

and 414). The temperature at which these crystals are in equilibrium with their mother liquor is the melting-point of pure ice when bathed in that liquor, and is lower than the melting-point of pure ice in contact with pure water. It is commonly called the freezing-point of the

Table 268.—Melting-points in Aqueous Solutions of Certain Gases

Only those gases for which the solubilities have been given in this compilation (Tables 230 and 231) are considered. The computed values of Δ have been derived from those solubilities at 0 °C by means of the relation $\Delta = 1.859N$. The experimental data, taken from the compilation by R. E. Hall and M. S. Sherrill,²²⁸ illustrate variations in the values of Δ/N when the assumptions that the solution is dilute and that there is no chemical reaction between the solute and solvent are not fulfilled.

F = formula-weight of the gas, N = number of gfw of gas dissolved in 1 kg of water, Δ °C = depression of the melting-point below what it would have been in the absence of the dissolved gas, the total pressure on the water being the same in both cases, P = partial pressure of the gas whenits saturated aqueous solution at 0 °C contains N gfw of gas per kg of water.

	Computed	Δ . $P = 1$ atm			- Experime	ntal Δ ·	
Gas	F	10 ⁻³ N	10 ⁸ ∆	Gas	N	Δ/N	Refa
Air	28.96	1.29	2.4 6	CO2	0.07	2.12	\mathbf{GF}
Α	39.91	2.4	4.3	$NH_{\bullet}OH$	0.006	2.11	J
He	4.00	0.43	0.8		0.01	2.03	Ĵ
\mathbf{Kr}	82.9	4.46	8.3		0.02	1.97	Ĵ
Ne	20.2	0.48	0.9		0.05	1.96	Ĵ
Rn	222.	22.8	42.	NH_a^c	0.5	1.86	R, P, S
Xe	130.2	10.8	20.1		2.0	1.9	R, P, S
$\mathbf{H}_{\mathbf{s}}$	2.015	0.96	1.8		5.0	2.0	R, P, S
N_2	28.02	1.03	1.9		10.0	2.2	R, P, S
Oz	32.00	2.18	4.0		15.0	2.5	R, P, S
O ₃	48.00	28.6	53.		20.0	2.9	R, P, S
CO	28.00	1.58	2.94		25.0	3.3	R, P, S
CO2	44.00	76.4	142.		30.0°	3.7	R, P, S
NH.	17.03	52600.f					,

^{*} References:

GF

ŀ

b If the formula-weight for air is taken as 30.00, the computed value of 1000Δ will be 2.3, as given by H. W. Foote and G. Leopold,²⁰⁰ and by H. W. Foote.²⁰⁰ This set of values is taken from p. 261 of Hall and Sherrill's report ²⁰¹; they give

another, and very slightly different, set on p. 255.

"Apparently the only available experimental value for the depression produced by CO is that by P. Falciola, and quoted in Hail and Sherrill's report (p. 255). He did not determine the concentration of the solution, has not reported the partial pressure of

Garelli, F., and Falciola, P., Gas. chim. ital., 34, II, 1-12 (1904).
Jones, H. C., Z. physik. Chem., 12, 623-656 (1893).
Postma, S., Rec. trav. chim. Pays-Bas, 39, 515-536 (1920).
Rupert, F. F., J. Am. Chem. Soc., 32, 748-749 (1910), superseding 31, 866-868 (1909).
Smits, A., and Postma, S., Proc. Akad. Wet. Amsterdam, 17, 182-191 (1914). R

²²⁸ Hall, R. E., and Sherrill, M. S., Int. Crit. Tables, 4, 254-264 (1928).

²²⁹ Foote, H. W., and Leopold, G., Am. J. Sci. (5), 11, 42-46 (1926).

²⁸⁰ Foote, H. W., Int. Crit. Tables, 4, 6 (1928).

²³¹ Hall and Sherrill, Int. Crit. Tables, 4, 254-264 (1928).

²⁸² Falciola, P., Gas. chim. ital., 39, I, 398-405 (1909).

Table 268.—(Continued)

the CO (it was probably about 1 atm), and gives but the single value $\Delta=0.015$ °C. As the solubility data indicate that 1000N=1.58 when P=1 atm, this value of Δ leads to $\Delta/N=9.5$, a very surprising value.

*Eutectic aqueous solution of NH₄OH; the eutectic point $(-\Delta)$ is -111 °C (Rupert places it below -120 °C; Postma, at -100.3 °C). According to Postma the composition of the eutectic is 34.5 per cent NH₃ by weight; i.e., N=30.9. At higher concentrations the solution consists of $(NH_4)_2O$ in NH_3OH ; and beyond that, of NH₄ in $(NH_4)_2O$ (Rupert) NH₈ in (NH₄)₂O (Rupert).

This is not an aqueous solution of NH3, but a dilute solution of water in NH3OH.

Table 269.—Melting-point in Sea-water

The data in terms of the salt-content (s) are from O. Krümmel 233 and are to be preferred to those in terms of σ_0 , by O. Pettersson.²³⁴ The latter are approximately represented by $t = 0.017 - \sigma_0/14.15$.

Salt content = s, density = $1 + 10^{-3}\sigma$; σ_0 , σ_t = value of σ at 0 °C, at t °C; melting-point (equilibrium between pure ice and sea-water) = t °C.

		Un	it of s	= 1 g sal	t per kg s	ea-wate	r; of den	sity = 1 g	g/ml		
s	-t	σ_l	\$	-t	σ_t	s	-t	σ_t	s	-t	$\sigma \iota$
1	0.055	0.72	11	0.587	8.80	21	1.129	16.87	31	1.683	24.96
2	0.108	1.52	12	0.640	9.60	22	1.184	17.67	32	1.740	25.76
3	0.161	2.34	13	0.694	10.41	23	1.239	18.49	33	1.797	26.58
4	0.214	3.15	14	0.748	11.22	24	1.294	19.29	34	1.853	27.39
5	0.267	3.96	15	0.802	12.02	25	1.349	20.10	35	1.910	28.21
6	0.320	4.75	16	0.856	12.84	26	1.405	20.91	36	1.967	29.02
7	0.373	5.57	17	0.910	13.64	27	1.460	21.71	37	2.024	29.83
8	0.427	6.38	18	0.965	14.45	28	1.516	22.52	38	2.081	30.65
9	0.480	7.19	19	1.019	15.25	29	1.572	23.34	39	2.138	31.46
10	0.534	8.00	20	1.074	16.07	30	1.627	24.14	40	2.196	32.27
σ_0		4.1	10).4	12.2		14.8	24	1.4	27.	1
	t	0.27	(0.715	0.85		1.025	1	.715	1.	895

solution, but the solution does not freeze as a unit unless it is saturated with the solute at the temperature at which freezing occurs. Under that condition, it freezes without change in concentration, and the resulting solid is an intimate mixture of solute and ice. A solution of that concentration is called a entectic solution.

If n is the number of effective gram-molecules contributed by the solute per kilogram of the solvent when the depression of the melting-point is Δ , we may write $\Delta = nE$, where E is characteristic of the solvent but independent of the nature of the solute, E varies with the concentration, and its limiting value (E_0) as n approaches zero is frequently called the cryoscopic constant of the solvent. By the number of "effective" molecules is meant the number of entities that individually participate in the thermal molecular agitation of the solution. If the solute is neither associated nor dissociated. and does not affect the molecular aggregation of the solvent, then $n = 1000 \times$ $m/FW \equiv N$, where m is the mass of the solute dissolved in the mass W of the solvent, and F is the formula-weight of the solute. If each molecule

²⁸⁸ Krümmel, O., "Handb. d. Ozeanographie," Vol. 1, 1907.

²⁸⁴ Pettersson, O., Vega-expedition; see Beibl. Ann. d. Physik, 7, 834-841 (1883).

of the solute is dissociated into two parts, and the solvent is not affected, then n = 2N; and similarly in other cases.

The quantity commonly tabulated is not E, but Δ/N , which equals E only when n = N and the state of molecular aggregation of the solvent is not affected by the presence of the solute.

The value of E is $RM_wT^2/1000L$, where R = gas-constant, $M_w = \text{molecular}$ weight of the vapor of the solvent, T = absolute temperature (273.1 + t) of the actual melting-point (t °C), and L = latent heat of fusion of the pure solvent at t °C; R and L must refer to the same unit of

Table 270.—Pressure-temperature Associations for Equilibrium Between the Several Pairs of Types of Ice

Adapted from the compilation a of P. W. Bridgman (ICT)²²¹ with addition of dP/dt from another paper.²²² (1912) See also Table 271 and Fig. 12.

P and t are the associated pressure and temperature, and dP/dt is the derivative under the condition that equilibrium is continually maintained.

Unit of P = 1 atm -1.0332 kg*/cm² = 1.0132 bars. Temp. = t °C dP/dtdP/dt-Ice-I and Ice-II--Icc-III and Ice-V-3358 2.66 1736 -75.0 8.95^{b} -35.08.95 1826 -65.03383 -30.02.66 -55.08.95 3395 -25.02.66 1916 3397 -24.3° 8.95 2006 -45.02.66 2094 -35.08.95 3409 -20.0-17.0°2100 -34.7° 3417 - Ice-V and Ice-VI -- Ice I and Ice III -2049 -- 60.0 + 5.2 6162 -20.00.77 -15.00.77 2091 -50.0+1.96166 2108 6169 -10.00.77 -40.0-0.66172 -5.00.77 2100 -34.7° 2087 6176 0.0 0.77 -30.0-3.1+0.16°2047 -22.0° 6175 -20.0-5.12035 -Ice-VI and Ice-VIId--Ice-II and Ice-III -19050 34.8 2100 -34.7°-80.02160 19720 -60.031.9 -34.0104 20320 28.6 2450 -31.0126 -40.020860 -20.024.2 2820 -28.0149 17.9 -25.0183 21290 0.0 3260 8,20 21530 +20.0 -24.3° 3397 3.38 21630 40.0 - Ice-II and Ice-V -0.97 21670 60.0 3397 -24.3° 21680 80.0 0 -25.0-65.23460 81.6° 21680 3680 -28.0-65.2-65.23880 -31.04070 -34.0

Based on P. W. Bridgman, 222 with a consideration of G. Tammann. 228

^o Triple point, see Table 263. ^d P. W. Bridgman ²²⁷; units changed by the compiler.

^b Bridgman (1912) gives $dP/dt = 8.35 \text{ kg}^*/\text{cm}^{2.\circ}\text{C} = 8.08 \text{ atm/}^{\circ}\text{C}$, which does not agree with the published values of P and t. Cf. note g to Table 272.

mass. For aqueous solutions so dilute that t may be taken as zero, L = 333.6 joules/g (value at 0 °C), T = 273.1 °K, R = 8.315 joules/gg-mole. K, and the cryoscopic constant is $E_0 = 8.315 M_w (273.1)^2 / 1000 \times$ $(333.6M_m) = 1.859$ °K per (g-mole per kg of water). Experimental data agree with this value as closely as should be expected. For more concentrated solutions, E takes other values, depending upon the melting-point.

Ice and Ice.

The known pressure-temperature associations for equilibrium between the members of the several pairs of ices are given in the preceding table.

PHASE DIAGRAM FOR WATER AND THE ICES

Although P. W. Bridgman and G. Tammann agree regarding the main features of the phase diagram for water and the several ices, they seem to

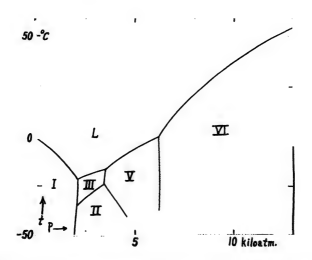


FIGURE 13. Phase Diagram for Water and the Ices (Bridgman).

[From Int. Crit. Tables, 4, 17 (1928).] The regions marked, I, II, III, V, and VI, are those in which the ices commonly designated by those symbols exist; L is the corresponding region for water. Bridgman has recently obtained evidence that an unstable form (IV) may arise within the region normally occupied by V, but cannot persist in the presence of V; and he has found a new stable form (VII) with triple-point L-VI-VII at 81.6 °C and 22400 kg*/cm². He has carried the pressure to 45000 kg*/cm² without obtaining evidence of any other form. See text.

Unit of P = 1000 atm = $1033.2 \, \text{kg*/cm}^2 = 1013.2 \, \text{megadyne/cm}^2$.

differ irreconcilably regarding the possible existence of the various unstable forms announced by Tanimann, and that introduces slight differences in the locations assigned by them for certain of the triple points and of the equilibrium curves in the neighborhood of those points. It seem's desirable for some one else to repeat the work of each, even though no evidence confirming the existence of Tammann's Ice-IV and other unstable forms has

been presented for many years.

Bridgman's complete diagram of 1912 is reproduced in Fig. 13. Since then he has extended his observations to 45,000 kg*/cm², finding another stable form (VII) at pressures beyond 22,400 kg*/cm² on the liquid-solid line,²³⁵ and an essentially unstable one, which he calls IV, that forms within the region normally occupied by V, but vanishes as soon as V appears.²³⁶

The corresponding numerical data given elsewhere in this compilation are based primarily on the work of Bridgman, for two reasons: (1) The program adopted for this compilation requires that the data be based as far as possible upon the *International Critical Tables*, and Bridgman's data are given in them. (2) The form in which his data have been presented is more readily adapted to this compilation than is that in which Tammann's data appear.

Those interested in the subject should consult the original articles by G. Tammann,²³⁷ by G. Tammann and E. Schwarzkopf,²³⁸ by G. Tammann and W. Jellinghaus,²³⁹ and by P. W. Bridgman.²²², ²³⁵, ²³⁶, ^{289a}

94. SURFACE CHARGES ON WATER AND ON ICE

At the interface between two substances there exists, in general, an electrical double-layer, the surface of one of the substances being charged positively and that of the other negatively; there is a contact difference of potential between the substances. Under the action of an impressed electrical field, the two components of this layer are urged in opposite directions, giving rise to the phenomena of cataphoresis and electric endosmosis. In many cases the components may be more or less separated by mechanical means also, giving rise to so-called frictional electricity and to the electrification produced by the spraying of a liquid in a gas, by the bubbling of a gas through a liquid, by the shattering of drops or streams of liquid (waterfall electricity), etc. It is with such phenomena that this section is concerned.

M. Faraday ^{239b} observed that when a stream of air containing minute drops of water flowed with pressure over surfaces of wood, brass, and certain other substances, those surfaces became negatively charged, but they did not become charged if the air was dry, even though the air contained uncondensed steam (¶2130, 2132). On the contrary, if the surface was of ice it became positively charged if the air was not dry (¶2131). These experiments are sometimes quoted as referring to the frictional electricity

²⁸⁵ Bridgman, P. W., J. Chem'l Phys., 5, 964-966 (1937).

²⁸⁶ Bridgman, P. W., Idem, 3, 597-605 (1935).

²⁸⁷ Tammann, G., Ann. d. Physik (4), 2, 1-31 (1900); "Kristallisieren und Schmelzen," 1903; Z. physik. Chem., 72, 609-631 (1910); 84, 257-292 (1913); 88, 57-62 (1914); "Aggregatzustände," 1922.

²³⁸ Tammann, G., and Schwarzkopf, E., Z. anory. allgem. Chem., 174, 216-224 (1928).

²³⁰ Tammann, G., and Jellinghaus, W., Idem, 174, 225-230 (1928).

²⁸⁰⁰ Bridgman, P. W., Proc. Am. Acad. Arts Sci., 48, 307-362 (1913); Z. physik. Chem., 86, 513-524 (1913); J. Franklin Inst., 177, 315-332 (1914).

²⁸⁹b Faraday, M., "Exp. Res. in Elec.," Vol. 2, pp. 106-126 (18th series, \$25), 1843.

generated by the rubbing of water on the solid, but it seems that the drops may have been shattered, which would have introduced complications. But F. Fairbrother and F. Wormwell ²⁴⁰ have reported that a rod of ice merely dipping into water becomes positively charged with respect to the water, whether the water is above or below 0 °C. This indicates that friction between ice and water will result in giving ice a positive charge, the same as Faraday observed in his experiments.

The interpretation of experiments on cataphoresis, electric endosmosis, electrification by bubbling, spraying, and shattering of drops and streams of liquid is difficult, there being many disturbing effects to be considered and eliminated. As a consequence, much of the subject is still in dispute, and it is not practical to give in this compilation more than a few rather general statements. Those interested should refer to the original papers, some of which are listed at the end of this section.

A mere rush of gas over the surface of a liquid gives rise to no charge unless the surface is ruptured. No charge arises from the simple condensation of a vapor nor from simple evaporation nor even from the violent boiling of water, not even though the surface were initially charged. That charge remains on the surface. When the interface between air and distilled water is suddenly ruptured, the net charge acquired by the air is always minus; that by the water is always plus (Gilbert and Shaw, 1925).* And J. J. Thomson (1894)* found that drops of water that have splashed from a wet plate are uncharged in water-vapor, charged positively in air and negatively in H_2 ; increasing the temperature increases the charge, the gas being air.

The water nuclei observed at the foot of a waterfall are negative if their diameters are under 80A ($1A = 10^{-8}$ cm), positive if between 80 and 150A, and uncharged if the diameters exceed 150A. To explain this, A. Bühl (1928)* has suggested that there are three electrified layers at an air-water surface: A negatively charged layer at the surface, a positively charged one about 80A below the surface, and a second negatively charged one about 150A below the surface. It is not clear how such a distribution can be maintained.

In her study of the cataphoresis of small particles in water, Miss Newton (1930)* observed that the particles (various oils and solids) always moved as if they were negatively charged, and that all of the same size had the same mobility if they were essentially spherical; irregularly shaped particles moved more slowly.

All these types of observation indicate that at an air-water interface the water is positively and the air negatively electrified, but estimates of the amount of the charge differ. It is, however, known that the amount of charge, and even its sign, may be greatly affected by the presence either of solutes or of surface impurities, and that it varies with the nature of the gas.

^{*} For complete reference see the following list of articles.

²⁴⁰ Fairbrother, F., and Wormwell, F., J. Chem. Soc. (London), 1928, 1991-1997 (1928).

The following articles will serve as an introduction to a study of these phenomena:

phenomena:
Alessandrini, E., Nuovo Cim. (5), 4, 389-402 (1902); Alty, T., Proc. Roy. Soc. London (A). 106, 315-340 (1924); Becker, A., Jahrb. d. Radioak., 9, 52-111 (1912) (bibliography of 60 entries); Bihl, A., Ann. d. Physik (4), 84, 211-244 (1927), 87, 875-908 (1928), Kolloid Z., 59, 346-353 (1932) (bibliography of 32 papers that have appeared since Becker's list); Busse, W., Ann. d. Physik (4), 76, 493-533 (1925); Currie, B. W., and Alty, T., Proc. Roy. Soc. London (A), 122, 622-633 (1929); Gilbert, H. W., and Shaw, P. E., Proc. Phys. Soc. London, 37, 195-214 (1925) (bibliography of about 100 references); v. Helmholtz, H., Ann. d. Physik (Wied.), 7, 337-382 (1879); Jurišić, P. J., Chem. Abs., 22, 1516 (1928) \(\infty \) Biochem. Z., 189, 294-301 (1927); Lachs, H., and Biczyk, J., Roczniki Chemii, 11, 362-375 (1931) (Polish, German summary); Lenard, P., Ann. d. Physik (Wied.), 46, 584-636 (1892), Sitz. Heidelberger Akad. Wiss. (A), 5, No. 28 (1914); Lignana, M., Atti d. R. Acc. Sci. Torino, 65, 276-281 (1930); McTaggart, H. A., Phil. Mag. (6), 27, 297-314 (1914); Mooney, M., Phys. Rev. (2), 23, 396-411 (1924); Nolan, J. J., Proc. Roy. Irish Acad., 37, 28-39 (1930); Nolan, P. J., Idem. 1, 417-428 (1926); Nolan, J. J., Proc. Roy. Irish Acad., 37, 28-39 (1926); Quincke, G., Ann. d. Physik (Pogg.), 113, 513-598 (1861); Reuss, F. F., Mém. de la soc. imp. naturalistes à Moscou, 2, 327-337 (1809) \(\rightarrow\) Wiedeman's "Die Lehre von der Electricität," 2 ed., Vol. 1, p. 993, 1893; Ruff, O., Niese, G., and Thomas, F., Ann. d. Physik (4), 83, 631-638 (1927); Simpson, G. C., Phil. Trans. (A), 209, 379-413 (1909); Thomson, J. J., Phil. Mag. (5), 37, 341-358 (1894); Zeleny, J., Phys. Rev. (2), 44, 837-842 (1933).

More recent articles:

Chalmers, J. A., and Pasquill, F., Phil. Mag. (7), 23, 88-96 (1937); Chapman, S., Physics, 5, 150-152 (1934), Phys. Rev. (2), 45, 135-136 (A) (1934), 49, 206 (A) (1936), 51, 145 (A) (1937), 52, 184-190 (1937), 53, 211 (A) (1938); Gilford, C. L. S., Phil. Mag. (7), 19, 853-878 (1935); Gostkowski, K., Acta Phys. Polon., 3, 343-345 (1934); Gott, J. P., Proc. Cambridge Phil. Soc., 31, 85-93 (1935); Kemp, I., Trans. Faraday Soc., 31, 1347-1357 (1935); Malarski, T., Acta Phys. Polon., 3, 43-74 (1934); Milhoud, A., Compt. rend., 198, 1586-1589 (1934), 200, 1091-1093 (1935); Mukherjee, J. N., Chaudhury, S. G., and Ghosh, B. N., Kolloid-Beih., 43, 417-463 (1936); Procopiu, S., Compt. rend., 202, 1371-1373 (1936); Terada, T., and Yamamoto, R., Proc. Imp. Acad. Japan, 11, 214-215 (1935).

IV. Phase Transition

95. Energy Changes Accompanying Phase Transition

Table 271.—External Work and Change in Volume during Phase Transition

 Δv^* = increase in specific volume when the transition takes place in the direction indicated by the arrow and at the indicated pressure (P) and temperature $(t \, ^{\circ}C)$ under which the two phases are in equilibrium.

The data for water → vapor are based on the values given in Tables 242, 250, and 255; the others have been adapted from P. W. Bridgman's compilation ¹ and other papers.² The data for the triple-points as given in the 1935 paper essentially agree with those in his earlier paper,³ which are the ones here given.

Unit of P=1 atm = 1.0132 bars; of $\Delta v^*=1$ cm³/g; of $P\Delta v^*=1$ joule/g. Temp. = t °C

			•		•		
t	P	Δv^*	$P\Delta v^*$	t	P	$\Delta \tau'^*$	$P\Delta v^*$
	Water	r → Vapor			Ice-I -	→ Ice-I1	
0	0.006027	+ 206290	+126.0	34.71	2100	-0.2178	- 46.31
100	1	1672.1	169.4	-35	2094	-0.2177	-46.19
200	15.347	126.04	196.0	-45	2006	-0.2170	44.10
250	39.26	48.81	194.2	55	1916	-0.2162	-41.97
300	84.79	20.24	173.9	-65	1826	-0.2154	-39.85
350	163.2	7.07	116.9	-75	1736	-0.2146	-37.74
	Wate	r → Ice-I			Ice-I -	Ice-III	
0	1	+0.0900a	+0.0091	-20	2035	-0.1773	-36.56
- 5	590	0.1016	6.07	-22b	2047	-0.1818	-37.70
- 10	1090	0.1122	12.39	-30	2087	-0.1919	-40.58
-15	1540	0.1218	19.01	-34.7 <i>t</i>	2100	-0.1963	-41.76
-20	1910	0.1313	25.41	-40	2108	-0.1992	-42.54
-22 b	2047	0.1352	28.04	-50	2091	-0.2023	-42.86
	Water	→ Ice-III		-60	2049	-0.2049	-42.53
-17.00	3417	-0.0241	- 8.34		Too TT	→ Ice-III	
-18.5	2820	-0.0301	- 8.60				
-20.0	2430	-0.0371	- 9.13	-24.30	3397	+0.0145	+ 4.99
-22.0b	2047	-0.0466	- 9.66	-25.0	3260	0.0148	4.89
	Wate	r → Ice-V		-28.0	2820	0.0164	4.69
0444			22.0	-31.0	2450	0.0179	4.44
0.16^{d}	6175	-0.0527	-32.9	-34.0	2160	0.0206	4.50
0.0	6160	-0.0527	- 32.89 32 .20	-34.77	2100	0.0215	4.58
5.0 10.0	5270 4360	-0.0603 -0.0679	-32.20 -29.99		Ton-TI	→ Ice-V	
-15.0	3680	-0.0754	-28.12	-24.30	3397	-0.0401	-13.80
-17.00	3417	-0.0788	-27.28	-25.0	3460	-0.0401	-14.05
-20.0	3040	-0.0828	-25.50	-28.0	3680	-0.0401	-14.95
20.0	3040	0.0020	20.00	-31.0	3880	-0.0401	-15 77
	Water	r → Ice-VI		-34.0	4070	-0.0401	-16.54
-15	4640	-0.0980	-46.1	•			2010
-10	5110	-0.0960	-49.8		Ice-III] → Ice-V	
- 5	5620	-0.0938	-53.4	-17.00	3417	-0.0547	-18.94
0	6160	-0.0916	-57.1	-20.0	3409	-0.05469	-18.89
+ 0.164	6175	-0.0916	-57.3	-24.30	3397	-0.0546	-18.79
5	6780	-0.0884	-60.7	-25.0	3395	-0.05461	-18.78
10	7390	-0.0844	-63.2	-30.0	3383	-0.05454	-18.68
15	8040	-0.0798	-65.0	-35.0	3358	-0.05446	-18.53

¹ Bridgman, P. W., Int. Crit. Tables, 4, 11 (1929).

² Bridgman, P. W., J. Phys'l Chem., 3, 597-605 (1935); 5, 964-966 (1937).

⁸ Bridgman, P. W., Proc. Am. Acad. Arts Sci., 47, 438-558 (1912).

			Table 2	271.—(Continuea)		
	P	Δv^*	$P\Delta v^*$	1	P	∆ y*	$P\Delta v^*$
	Wat	er → Ice = VI-			Ice-V -	→ Ice-VI	
20	8710	-0.0751	-66.3	+ 0.16d	6175	-0.0389	-24.34
30	10250	-0.0663	-68.9	0.0	6176	-0.03886	-24.32
40	11990	-0.0590	-71.7	- 5.0	6172	-0.03866	-24.07
50	13970	-0.0523	-74.0	-10.0	6169	-0.03847	-23.94
60	16150	-0.0477	-78.0	-15.0	6166	-0.03828	-23.81
•		r → Ice-VI		-20.0	6162	0.03809	-23.68
				`	Ice-VI	→ Ice-VII	
52.5	14518	-0.0508	-74.7	-80.0	19047		
57.2	15485	-0.0478	-74.9	-60.0	19715		
66.0	17421	-0.0424	-74.7	-40.0	20325		
73.8 80.8	19357 21292	-0.0376 -0.0335	-73.8 -72.3	-20.0	20857		
81.6	21680	-0.0333	-72.4	0.0	21292	-0.0567	-122.6
81.0-			14.7	+20.0	21534	-0.0570	-124.5
		r → Ice-VII —		40.0	21631	-0.0573	-125.5
81.6*	21680	-0.0910	200.0	60.0	21670	-0.0576	-126.5
95.3	23228	-0.0879	-206.9	80.0	21680	-0.0580	-127.5
110.3	25164	-0.0847	-215.7				
124.1	27100	-0.0817	224.6	81.6	21680	-0.0580	-127.5
137.1	29035	-0.0789	-231.4				
149.5	30971	-0.0763	-239.3				
161.1	32906	-0.0738	-246.1				
172.1	34842	-0.0715	-253.0				

Table 271 (Continued)

^a See also Table 201.

182.5 192.3

- Triple-point of water, ice-I, and ice-III.
 Triple-point of water, ice-III, and ice-V.
 Triple-point of water, ice-V, and ice-VI.
 Triple-point of water, ice-VI, and ice-VII.
 Triple-point of ice-I, ice-II, and ice-III.

- "Triple-point of ice-II, ice-III, and ice-V.

Latent Heat of Phase Transition.

By the latent heat (L) involved in a given change in phase, is meant the amount of heat absorbed per unit of mass isothermally and reversibly transformed in the indicated direction. The two phases are assumed to be and to remain in mutual equilibrium.

Latent Heat of Vaporization.—For a review of the several determinations of the latent heat of vaporization of water prior to 1930, see E. F. Fiock.⁴ Osborne and his associates have redetermined the latent heat from 50 °C almost to the critical point, and have concluded that the best formulation for $t \equiv 100$ °C is that of expression (1),

$$L = 1585.19 \left(\frac{374.15 - t}{100} \right)^{0.404} - 36.75304 \left(\frac{310 - t}{100} \right)^{1.73} + 17.9218 \times \left(\frac{165 - t}{100} \right)^{2.2}$$
 (1)

the unit being 1 Int.joule/g, and temperature being t °C on the international scale.⁵ This supersedes previous formulations and tables of L published by these workers, including that of Fiock,4 Osborne, Stimson, and Fiock, Fiock and Ginnings, and Osborne, Stimson, and Ginnings.8

⁴ Fiock, E. F., Bur. Stand. J. Res., 5, 481-505 (RP210) (1930).

⁵ Osborne, N. S., Stimson, H. F., and Ginnings, D. C., J. Res. Nat. Bur. Stand., 18, 389-448 (RP983) (1937).

Their work, however, is continuing at the two extremes of the temperature range. As a result, the extreme values given by (1) are subject to revision; an idea of the size of the changes that may be expected can be obtained from Table 272, containing the most recent revision. Another recent formulation, differing from (1), has been proposed by M. Jakob and W. Fritz⁹; the values defined by it are indicated in the JF column of Table 272.

Of the older data given in Table 272 for comparison, those of H. L. Callendar are unique in that they are based upon a formulation that assumes that identity between liquid and vapor does not occur until the temperature reaches 380.5 °C, 6.5 °C above the critical temperature at which the meniscus vanishes.10

A. W. Smith 11 has reported that the value found for the latent heat of vaporization when determined from a slow evaporation from a still surface is about 0.75 per cent greater than when it is determined from the evaporation produced by actual boiling. He thought that the difference was probably real. If it is, then it indicates that the vapor as it leaves the liquid is polymeric, but quickly breaks down into ordinary water-vapor (cf. the next paragraph, Sublimation).

Latent Heat of Sublimation.—Previous to the observations of H. T. Barnes and W. S. Vipond 12 it was assumed that the latent heat of sublimation of ice-I at 0 °C was the sum of the latent heat of fusion of ice-I at 0 °C and the latent heat of vaporization of water at the same temperature; and the same is quite generally assumed today. This assumption implies that the vapor in immediate contact with ice is identical with that which is in equilibrium with water, which second assumption need not be In fact, Barnes and Vipond reported it false. They stated that when vapor is removed very quickly from ice, the latent heat is only 2540 joules/g, whereas if it is removed slowly the latent heat is 2930 joules/g. The first is only 46 joules/g (=1.6%) greater than the latent heat of vaporization of water; whereas the second exceeds the sum of the latent heats by 102 joules/g (=3.5%). These figures, of which the compiler has found no confirmation, indicate that the vapor as it leaves the ice is polymeric, and only later breaks down into ordinary water vapor, absorbing heat in the process. Barnes thought that it was probable that in ordinarv evaporation from ice and snow the change of the vapor from the solid into ordinary vapor takes place just outside the surface, and under ordinary

⁶ Osborne, N. S., Stimson, H. F., and Fiock, E. F., Bur. Stand. J. Res., 5, 411-480 (RP209) (1930) = Trans. Am. Soc. Mech. Eng., 52, 191-220 (FSP-52-28) (1930).

⁷ Fiock, E. F., and Ginnings, D. C., Idem, 8, 321-324 (RP416) (1932).

⁸ Osborne, N. S., Stimson, H. F., and Ginnings, D. C., Mech. Eng., 56, 94-95 (1934); 57, 162-163 (1935).

⁹ Jakob, M., and Fritz, W., Physik Z., 36, 651-659 (1935). Also Jakob, M., Mech. Eng., 58, 643-660 (1936).

 ¹⁰ For his reasons, see Callendar, H. L., Proc. Roy. Soc. London (A), 120, 460-472 (1928);
 Proc. Inst. Mechan. Eng. 1529, 507-527 (1929). See also p. 558+.
 11 Smith, A. W., J. Opt. Soc. Amer., 10, 711-722 (1925).

¹² Barnes, H. T., and Vipond, W. S., Phys. Rev., 28, 453 (A) (1909).

circumstances the difference between the two would escape detection in vapor pressure measurements.13

Latent Heat of Fusion.—The determinations prior to 1871 of the latent heat of fusion of ice at 0 °C have been reviewed by A. W. Smith, 14 those between 1870 and 1913 by H. C. Dickinson, D. R. Harper, and N. S. Osborne 15; and recently the entire subject has been again reviewed by A. W. Smith.¹¹ In the second paper, Smith's table summarizing the early determinations is republished; those values need not detain us. Four values between 1870 and 1913 have to be considered—those obtained by A. W. Smith, 14 by A. D. Bogojawlensky, 16 by U. Behn, 17 and by C. Dieterici.¹⁸ These, after correction of the last two by W. A. Roth, ¹⁹ and the reduction of all to the same basis by Dickinson, Harper, and Osborne, lie between 79.59 and 79.69, and their mean is 79.62 cal₁₅/g (=333.21 joule/g) at 0 °C. In the same paper, Dickinson, Harper, and Osborne publish the results of a very careful determination by themselves, embracing 21 sets of observations and referring to ice from several sources. Their results lie between 79.57 and 79.68, mean = $79.63 \text{ cal}_{15}/g$, which, when corrected for the use of a slightly erroneous value for the specific heat of ice,20 becomes 79.69 cal₁₅/g (= 333.66 joule/g)* at 0 °C. Additional very careful determinations by Dickinson and Osborne 20 gave 333.63 joule/g at 0 °C. More recently O. Maass and L. J. Waldbauer, 21 and O. Maass and W. H. Barnes ^{13, p. 30} have obtained the lower values † 79.42 and 79.40 cal/g, about 332.5 joule/g; the particular calorie used is not stated, and the researches appear to have been much less elaborate than those carried out at the Bureau of Standards. The value obtained by that Bureau and published in the International Critical Tables is given in Table 272.

As long ago as 1850, C. C. Person 23 pointed out that there is an incipient melting of ice before the temperature has risen to the true melting point, and that to ignore this may seriously affect the value obtained for the latent heat of fusion. J. Y. Buchanan 24 called attention to the same thing, and showed that such melting arises from the presence of impurities

^{*} The energy was measured electrically and reduced to cal₁₈ on the assumption that 1 cal₁₅ = 4.187 joules. This values, which is greater than the one (4.185) now accepted, was used in the reverse conversion.

[†] These seem to have been derived from the work of W. H. Barnes and O. Maass 22: see remarks in Section 69.

¹⁸ Barnes, H. T., "Ice Engineering," p. 33, 1928.
¹⁴ Smith, A. W., Phys. Rev., 17, 193-232 (1903).

¹⁶ Dickinson, H. C., and Harper, D. R., and Osborne, N. S., Bull. Bur. Stds., 10, 235-266 (S209) (1913).

¹⁶ Bogojawlensky, A. D., Schrift. Dorpater Naturf. Ges., 13, (1904).

¹⁷ Behn, U., Ann. d. Physik (4), 16, 653-666 (1905).

¹⁸ Dieterici, C., Idem, 16, 593-620 (1905).

¹⁰ Roth, W. A., Z. physik. Chem., 63, 441-446 (1908).

²⁰ See Dickinson, H. C., and Osborne, N. S., Bull. Bur. Stds., 12, 49-81 (S248) (1915).

²¹ Maass, O., and Waldbauer, L. J., J. Am. Chem. Soc., 47, 1-9 (1925).

²² Barnes, W. H., and Maass, O., Can. J. Res., 3, 70-79, 205-213 (1930).

²⁸ Person, C. C., Ann. Chim. et Phys. (3), 30, 73-81 (1850).
24 Buchanan, J. Y., Proc. Roy. Inst'n Grt. Brit., 19, 243-276 (1908); Proc. Roy. Soc. Edinburgh, 14, 129-149 (1887) → Nature, 35, 608-611; 36, 9-12 (1887).

Table 272.—Latent Heat of Change in Phase

The latent heat (L) is the amount of heat absorbed per unit of mass isothermally and reversibly transformed in the direction indicated by the arrow; $\Delta v^* =$ increase in the specific volume associated with that transformation; P = pressure at which the two phases are in equilibrium at the indicated temperature; $\Delta E = L - P\Delta v^* =$ increase in internal energy, and L/T = increase in entropy, each per unit of mass transformed. The values of $P \cdot \Delta v^*$ have been taken from Table 271.

For the water \rightarrow vapor data the O values are to be preferred. They are given directly The values corresponding to any of the other sets of data will be obtained by adding the appropriate value of ΔL to the corresponding O value of L; e.g., the KSG value for 10 °C is 2476 48 - 0 30 = 2476.18; the WT value for 0 °C is 2500.00 - 11 = 2489.

Unit of L, $P \cdot \Delta v^*$, and $\Delta E = 1$ Int joule/g = 0.2389 cal₁/g Temp = $t \circ C = (273.1 + t) \circ K$ I. Water \rightarrow Vapor

1.	water ->	v apoi							
Refa→	o L	OSG	KSG	Łck	Koch ΔL	WT	JΓ	С	ICT
0 10 20 30 40	2500 00 2476 48 2452 93 2429 30 2405 54		+0 15 -0 30 -0 51 -0 42 -0 40			-11 - 7 - 5 - 3 - 2	- 39 - 25 - 15 - 10 - 03		- 6 - 5 - 4 - 4
50 60 70 80 90	2381 58 2357 37 2332 83 2307 87 2282 42		-0 21 +0.03 +0 22 +0 31 +0 33			0 0 0	- 02 + 02 + 01 - 05 - 10		- 2 0 0 + 1 + 2
100 110 120 130 140	2256 37 2229 64 2202 13 2173 67 2144 22	+0 41 +0 23 +0 07 -0 02	+0 27 +0 16 +0 05 +0 01 -0 03			0 - 1 - 2 - 4 - 4	- 1 4 - 2 2 - 2 7 - 3 2 - 3 9		+ 2 - 3 - 4 - 6
150 160 170 180 190	2113 74 2082 12 2049 17 2014 61 1978 28	+0 02 -0 02 +0 01 0	-0 14 -0 34 -0 56 -0 66 -0 65			- 4 - 2 + 1 + 5	- 43 - 51 - 56 - 57 - 54		- 7 - 8 -10 -15
200 210 220 230 240	1940 08 1899 84 1857 41 1812 58 1765 13	-0 01 -0 02 +0 02 0	-0 56 -0 84 -0 31 -0 21 -0 18				- 53 - 48 - 42 - 34 - 24	+10 1 +11 9 +12 8 +14 9 +15 5	
250 260 270 280 290	1714 77 1661 17 1603 92 1542 47 1476 17	0 +0 02 +0 01 +0 01 0	-0 25 -0 43 -0 74 -1 13 -1 60				- 19 - 10 - 02 + 05 + 10	+16 9 +19 0 +20 1 +20 9 +22 7	
300 310 320 330 340	1404 09 1324 92 1237 24 1139 29 1027 00	-0 01 +0 01 -0 01 -0 01 0	-2 01 -2 12 -2 02 -2 15 -2 01		-2 2 -2 9 -2 3 -2 0 -1 4		+ 20 + 29 + 40 + 40 + 40	+23 6 +24 5 +25 1 +25 0 +23 5	
350 355 360 365 366	892 85 812 98 719 43 603 25 575 69	-0 01 -0 02 +0 02	-0 70 +5 30	+ 20 + 12 + 23 + 24	-08 -01 -02 +04		+ 30 + 27 + 38	+22 0 +20 0	
367 368 369 370 371	546 04 513 79 478 24 438 30 392 10	-0 01 +0 01 +0 02 +0 02 +0 13		+ 58	+04		+ 88	+39 0	
372 373 374 374.15	336 04 260 98 114 61 0	+0 10 -0 19 +0 09 0		+13 1 +54 5 0	-0 3		+13 5	e+188 1 ²	

Table 272.—(Continued)

II. Ice → Vapor (See text, Sublimation.)

III. Water → Ice; Ice → Ice. P. W. Bridgman, a

t	L	$P \cdot \Delta v^*$	ΔE	L/T	<i>t</i>	L	$P \cdot \Delta v^*$	ΔE	L/T
		Vater → I				Ice-	·I → Ice-II	7	
o_	333.66	+0.0091	-333.6	-1.22	-34.7h	-42.3	-46.3	4.0	-0.177
- 5 10	-308. ₅	6.1	-314. ₈	-1.15	-35	-42.48	-46.19	3.71	-0.178
-15	-284.8 -261.6	12.4 19.0	- 297.2 - 280.6	-1.08 -1.01	-45	-40.50	-44.10	3.60	-0.177
-20	-241.4	25,4	-266. ₈	-0.95	-55 -65	-38.59 -36.70	-41.97 -39.85	3.38	-0.177 -0.177
-220	-234.8	28.0	-262.8	-0.93	-75	-30.70 -34.77	-37.74	3.15 2.97	-0.177 -0.175
		ater → Ice					·I → Ice-II		0.175
-17.0^{d}	-257	- 8.3	-249	-1.004	-20	23.4	-36.6	60.0	0.092
-18.5	-240	- 8.6	-231	-0.94_3	-220	21.8	-37.7	59.5	0.087
-20.0	-226	- 9.1	-217	-0.89_3	-30	14.6	-40.6	55.2	0.060
-22.0°	-213	- 9.7	-203	-0.84_{8}	-34.74	9.2	-41.8	51.0	0.039
-		/ater → Ic			-40 -50	2.9	-42.5	45.4	0.012
+ 0.16* 0.0	- 293.4 293	-33.0 -33	-260.4 -260	-1.07_4	-50 -60	- 8.8 -23.0	-42.9 -42.5	34.1 19.5	-0.039 -0.108
- 5.0	-295 -285	-33 -32	-253 -253	-1.07_3 -1.06_8	00		II → Ice-II		-0.108
-10.0	-276	-30	-246	-1.04 _p	-24.31				0.004
-15.0	-265	-28	-237	-1.02_{0}	-25.0	70.7 68.2	5.0 4.9	65.7 63.3	0.284 0.2 75
-17.0d	-261	-27	-234	-1.026	-28.0	60.7	4.7	56.0	0.248
-20.0	-253	-26	-227	-1.00_{0}	-31.0	55.2	4.4	50.8	0.248 0.228
	W	ater - Ice	e-VI		-34.0	51.9	4.5	47.4	0.221
-15.0	-247	-46	-201	-0.95_{7}	-34.7h	51.5	4.6	46.9	0.216
-10.0	-264	-50	-214	-1.004			·II → Ice-V		
- 5.0	-281	-53	-228	-1.049	-24.34	67.0	-13.8	80.8	0.269
0 + 0.16*	-295 -294	-57 -57	-238 -237	-1.08_0 -1.07_5	-25.0 -28.0	67.0 66.1	-14.0 -15.0	81.0	0.270 0.270
5.0	-303	-61	-242	-1.09_0	-31.0	65,2	-15.8	81.1 81.0	0.269
10.0	-311	-63	-248	-1.09 ₀	-34.0	64.4	-16.5	80.9	0.269
15.0	-316	-65	-251	-1.09_{7}		Ice-	III → Ice-	V	
20 0	-320	-66	-254	-1.09_1	-17.0d	-3.7	-18.94	15.1	-0.014
30.0	-330	-69	-261	-1.08_{9}	-20.0	-3.72	-18.89	15.17	-0.014_{7}
40.0	-342	-72	-270	-1.09_{2}	-24.3	-3.7	-18.79	15.0	-0.015_{2}
50.0 60.0	-357 -379	-74 -78	-283 -301	-1.105	-25.0 -30.0	-3.64 -3.56	-18.78	15.14	-0.0147
00.0		ater → Ic		-1.13_{8}	-35.0	-3.50 3.47	-18.68 -18.53	15.12 15.06	-0.014 ₆ -0.014 ₆
F2.F				1.024			-V → Ice-V		0.02 %
52.5 57.2	-333.5 -336.5	-74.7 -74.9	-258.8 -261.6	-1.024 -1.019	+ 0 16*	-0.8	-24.34	23.5	-0.0031
66.0	-339.1	-74.7	-264.4	-1.009	0.0	-0.83	-24.32	23.49	-0.0030
73.8	-345.3	-73.8	-271.5	-0.995	- 5.0	-0.82	-24.07	23.25	-0.0031
80.8	-352.0	-72.3	-279.7	-0.995	-10.0	-0.80	-23.94	23.14	-0.0030
81.6	-354.5	-72.4	-282.1	-0.999	-15.0	-0.77	-23.81	23.04	-0.0030
		ater \rightarrow Ice			-20.0	-0.76	-23.68 VI → Ice-V	22.92	-0.0030
81.6 <i>f</i> 95.3	-354.5 -398.0	-200 0	-154.5 -191.1	0.999 1.080	0.0	-23.8	V1 → 1ce-V -122.6	+ 98.8	-0.087
110.3	398.0 444.4	-206.9 -215.7	-228.7	-1.159	20 0	-13.8	-124.5	+110.7	-0.047
124.1	-474.6	-224.6	-250.0	-1.195	40.0	- 6.3	-125.5	+119.2	-0.0048
137.1	-500.1	-231.4	-268.7	-1.219	60.0	- 2.1	-126.5	+124.4	-0.0015
149,5	-526.9	-239.3	-287.6	-1.247	80.0	0	-127.5	+127.5	0
161.1	-554.1	-246.1	-308.0	-1.276	81.67	0	-127.5	+127.5	0
172.1	-582.6	-253.0	-329.6	-1.309					
182.5 192.3	-610.2 -642.4	-258.9 -264.8	-351.3 -377.6	-1.339 -1.380					
476.07	074.7	204.0	311.0	1.000					

* References:

- Bridgman, P. W.^{2, 8} From the 1937 paper come the second set of data for water to ice-VI and all for ice-VII. The triple-point data given in the 1935 paper agree essentially with those in the 1912 one. Callendar, H. L., *Proc. Inst. Mech. Eng.*, 1929, 507-527 (1929).
- Eck, H., Tätigkeit Phys.-Techn. Reichs. im 1936, p. 32 = Physik. Z., 38, 256 (1937). Eck Compilation by Smith, A. W., and Bridgeman, O. C., Int. Crit. Tables, 5, 138 (1929).
- ICT Jakob, M., and Fritz, W., Physik. Z., 36, 651-659 (1935). Supersedes similar data by Jakob, M., Wiss. Abh. Phys.-Techn. Reichs., 12, 435-446 (1928) = Forsch.-Arb. Gebiete Ingenicuru., 310, 9-19 (1928) and by Jakob and Fritz. Wiss. Abh. Phys.-Techn. Reichs., 13, 93-111 (1928) = Z. Ver. deuts. Ing., 73, 629-636 (1929); Tech. Mech. Thermodynam., 1, 173-183, 236-240 (1930); Forsch. Gebicte Ingenieuru., 4, 1905 (1932) JF 295-299 (1933)
- Koch, W., Idem, 5, 257-259 (1934) → Z. Ver. deuts. Ing., 78, 1160 (1934). Koch
- Keyes, F. G., Smith, L. B., and Gerry, H. T., Proc. Am. Acad. Arts Sci., 70, 319-KSG 364 (1935)

Table 272.—(Continued)

- O Osborne, N. S., Private communication, 1938.
- OSG Osborne, N. S., Stimson, H. F., and Ginnings, D. C., superseding both Mech. Eng., 57, 162-163 (1935) and Osborne, Stimson, and Fiock.
- WT Holborn, L., Scheel, K., and Henning, F., "Warmetabellen," 1919.
- ^b This latent heat of fusion of ice-I is from the compilation by R. de Forcrand and L. Gay ^{2b} and is based on observations by H. C. Dickinson and N. S. Osborne.^{2b}
 - "Triple-point of water, ice-I, and ice-III.
 - d Triple-point of water, ice-III, and ice-V.
 - * Triple-point of water, ice-V, and ice-VI.
 - Triple-point of water, ice-VI, and ice-VII.
- "In computing these values of L for ice-I to ice-II, Bridgman used dP/dt = 8.08 atm/°C instead of the value (8.95) defined by the values of P in Table 270, presumably in order to make the L's satisfy the triple-point condition. He used the same value for all temperatures. It is obvious that the values given for -34.7 °C are discordant with those given for lower temperatures.
 - * Triple-point of ice-I, ice-II, ice-III.
 - 'Triple-point of ice-II, ice-III, and ice-V.
- *The values of ΔH given in Bridgman's Table XXI are obviously inconsistent with the values of Δv and of the adjusted values of dP/dt given in the same table. The values of L here given have been computed by the compiler from those values of Δv and adjusted dP/dt, and the values of ΔE have been changed accordingly.
- ¹ Callendar gives the following values for the latent heat of vaporization at higher temperatures: 375 °C, 245.3; 377 °C, 186.3; 380 °C, 48.1; and 380.5 °C, 0 joule/gram.

included in the ice. Nevertheless, this source of error has frequently been overlooked. It has, however, been carefully considered in the more elaborate of the recent determinations.

Latent Heat of Ice to Ice.—The values given in Table 272 for the latent heat involved in the transition of one form of ice to another are based almost exclusively upon the work of P. W. Bridgman ²⁵ and have been computed by means of Clapeyron's equation: $L = T \cdot \Delta v^* \cdot dP/dT$, where $T \circ K \ (\equiv 273.1 + t \circ C)$ is the absolute temperature, and Δv^* is the increase in specific volume when the transition takes place in the indicated direction.

Miscellanca.—P. Walden ²⁶ has announced that ML/T has nearly the same value (13.5 cal/°K per g-mole = 56.5 joule/°K per g-mole) for all normal liquids when they freeze; i.e., if the molecular weight (M) of the liquid is 18.015 the increase in entropy during transition is L/T = 3.14 joule per gram. °K. Values of L/T are included in Table 272.

From the fact that L/T has, roughly, the same value whether water freezes to ice-I, ice-III, ice-V, or ice-VI, G. Tammann concluded that these four types of ice are isometric, differing only in the distance between adjacent molecules, not in the grouping of atoms in the molecules.²⁷

- 25 Bridgman, P. W., Proc. Am. Acad. Arts Sci., 47, 439-558 (1912).
- 26 Walden, P., Z. Elektrochem., 14, 713-724 (1908).
- 27 See Tammann, G., "Agregatzustände," p. 143-144, 1922.
- 28 de Forcrand, R., and Gay, L., Int. Crit. Tables, 5, 131 (1929).
- Dickinson, H. C., and Osborne, N. S., Bull. Bur. Stand., 12, 49-81 (S248) (1915).

Disposable Energy from Isopiestic Change in Phase.

The maximum amount of external work that can be obtained from an isothermal change in phase under a constant pressure is $W_{TP} + P \cdot \Delta v$, where Δv is the increase in volume. As an amount of work equal to $P \cdot \Delta v$ must be expended against the pressure P, only the amount W_{TP} is disposable for other purposes. Proceeding as in Section 6, and using the same constants as were used there, one obtains formulas (2), (3), and (4),

Table 273.—Disposable Energy from Isopiestic Change in Phase

When the change occurs isothermally and at a constant pressure, the disposable energy is $W_{TP} = w - f(T, P)$; $f(T, P) = +0.008315T \log_{\theta}(P/A)$ for each of the changes water \rightarrow vapor and ice-I \rightarrow vapor, and

$$f(T, P) = + \int_{A}^{P} (\Delta v)_T dp$$
 for water \rightarrow ice-I. The following values have

been computed by means of formulas (2), (3), and (4); $w_g = w/18.0154$. For the change water \rightarrow vapor, w is exactly zero at 100 °C; the finite value (-0.04) defined by formula (2) arises from errors in the constants used in deriving the formula.

Unit of $w = 1 \text{ kj/gfw-H}_2O$; of $w_g = 1 \text{ j/g}$. Temp. = $t \circ C = T \circ K$

Cha	Change →		water → vapor		vapor	water → ice-I		
t	T	76'	v_{g}	w	70° g	7 0	w_g	
-30	243.1	-15.33	-851	-15.92	-882	+0.587	+32.6	
-20	253.1	-14.07	-781	-14.48	804	+0.413	+22.8	
-10	263.1	-12.83	-712	-13.04	-724	+0.213	+11.8	
0	273.1	- 11.60	-644	-11.60	644	0	0	
+10	283.1	-10.39	-577					
20	293.1	-9.18	-510					
40	313.1	-6.82	-379					
60	333.1	-4.51	-250					
80	353.1	-2.24	-125					
100	373.1	-0.04	-2					

which are indeed merely the differences between the equations there obtained for the corresponding pairs of phases, excepting the last term in (4), which was there ignored as negligible with respect to the large value of W for the formation of the individual phase. As before, A denotes the pressure of 1 atm, temperature = t °C = T °K. Values computed by means of those equations are given in Table 273.

Water to vapor:

$$(W_{TP}) = -11.599 + 0.12216t - 2.697(t/1000)^2 + 1.5473(t/1000)^3 - 10.517 \times \{(T/273.1)\log_e(T/273.1) - t/273.1\} - 0.008315T\log_e(P/A)$$
kj/gfw (2)

Ice-I to vapor:

$$(W_{TP}) = -11.599 + 0.14417t - 71.622(t/1000)^{2} + 1.5473(t/1000)^{3} + 9.941 \times \{(T/273.1)\log_{e}(T/273.1) - t/273.1\} - 0.008315T\log_{e}(P/A)$$
kj/gfw (3)

Water to ice-I:

$$(W_{TP}) = -0.02201t + 68.925(t/1000)^2 - 20.458\{(T/273.1) \log(T/273.1) - t/273.1\} - \int_A^P (\Delta v)_T dp \text{ kj/gfw}$$
(4)

96. VAPORIZATION AND CONDENSATION

With certain restrictions which will appear, this section may be said to deal exclusively with kinetic phenomena. In it are considered those phenomena that accompany the transition, direct or reverse, between watervapor and a condensed phase—water or ice—and that cannot be derived directly and solely from observations made under equilibrium conditions. Among the data that are thus excluded are: pressure-volume-temperature associations for (a) dilated water-vapor, Section 14, (b) water and steam at saturation, Section 88, (c) ice and saturated vapor, Section 92; steamtables and diagrams, Section 90; energy changes accompanying phase transition, Section 95.

Two distinct classes of problems have to be distinguished. One is concerned with the escape of molecules from the denser phase, and with their capture by it. The other has to do with the net transfer of substance from one phase to another under certain specific conditions—with the lack of balance between the escape and the capture of molecules under those adventitious conditions. Problems of the first class are the more fundamental; those of the second are of the greater technical importance, and to them the terms "evaporation" and "condensation" will be restricted, in accordance with common usage.

Escape and Capture of Molecules.

General Relations.—Let α = the ratio of the number of vapor molecules that are caught by the surface of the denser phase (liquid or solid) to the number that strike it in the same time, and m_s = total mass of the vapor molecules that strike unit area of the surface in unit time when the vapor is saturated with reference to the surface. Then the total mass of those vapor molecules that enter the denser phase, per unit area and per unit time, under those conditions will be αm_s , and the corresponding number, m_e , that escape will be the same, $m_e = \alpha m_s$. All three quantities depend upon the temperature of the surface itself.

The quantity α , here called coefficient of capture, is sometimes called the accommodation coefficient, and the failure of the surface to catch all the molecules striking it is sometimes described as a reflection of the molecules.

From a knowledge of the temperature and of the molecular weight and pressure of the saturated vapor, m_s can be readily computed within a small range of error, depending upon the departure of the vapor from the ideal state, and that error can be allowed for whenever other conditions justify the trouble. Hence α can be determined if m_e can be measured. Such

is the procedure followed, it being assumed that m_e and α are each independent of the pressure of the overlying vapor, and that the vapor and the surface are at the same temperature. Unless the radius of curvature of the surface is great with reference to the mean free path of the molecules of vapor, m_e and α may be expected to depend upon that curvature. In the rest of this section it will be assumed that the surface is essentially plane.

If n = the number of molecules per unit volume of the saturated vapor and $\bar{v} =$ their mean translational velocity, then the number (n_s) that strike one side of a unit area per unit of time is* $n_s = n\bar{v}/4 = NP_{\rm sat}/(2\pi MRT)^{0.5}$, and $m_s = n_s M/N$ (see Table 12). For water, $10^{-18}n_s = 6.24 P_{\rm sat}/T^{0.5}$ molecules per cm²sec, and $m_s = 0.1857P_{\rm sat}/T^{0.5}$ mg per cm²sec, the unit of $P_{\rm sat}$ being 1 dyne/cm², and the temperature being T °K.

The determination of m_e is exceedingly difficult, resting upon extrapolation from observed rates of evaporation that are less than 0.005 and often less than 0.00002 as great as m_e , and as yet, only inferior limits to m_e have been obtained. The observed rate of evaporation is limited by (1) the rate at which the vapor can be removed from the surface of the denser phase, and (2) the rate at which heat can be supplied to the surface (see Table 274). The first is very seriously limited by the presence of a stagnant layer of gas (or vapor) that always clings to the surface, and through which the vapor passes by the slow process of diffusion (see p. 624).

That m_e is finite, that at a given temperature there is a definite limit to the rate of evaporation of a given liquid, has long been recognized.⁸⁰

Coefficient of Capture.—The only available estimates of α for water seem to be those of T. Alty, 31 of T. Alty and C. A. Mackay, 32 and of T. Alty and F. H. Nicoll, 33 of which that of 1935 supersedes all the others. In that, the temperature of the surface is inferred from the observed surface tension, and the value found for α is 0.036 for a surface temperature of 10 °C. The corresponding value of m_e is 4.9 mg/sec·cm², which represents a thermal current of 12.13 watts/cm² through the surface, which in turn could be supplied by conduction through the water only if the temperature gradient were 2100 °C/cm. Obviously, convection must play a most important part in supplying the necessary heat. Their earlier and less accurate work (Alty, 1931; Alty and Nicoll, 1931) indicated that the value

^{*} If all the quantities are expressed in cgs units, then the unit of n_s will be 1 molecule/cm²sec; if all except P_{nnt} are in such units and the unit of P is q dynes/cm², then the unit of n_s will be q molecules/cm²sec. The quantity RT is of the nature of pressure times specific volume; if the unit of this pressure as well as that of P_{nnt} is q dynes/cm², those of mass and volume being 1 g-mole and 1 cm³, respectively, then that of n_s will be $q^{0.5}$ molecules/cm²sec. If the unit of pressure is 1 mm-Hg, q = 1333.22 and $q^{0.5} = 36.513$.

³⁰ See Mache, H., Sitzb. Akad. Wiss. Wicn (Abt. IIa), 119, 1399-1423 (1910); Z. Physik, 107, 310-321 (1937), and for earlier observations related to this subject, Winklemann, A., Ann. d. Physik (Wied.), 22, 1-31 (1884); 23, 203-227 (1884); 26, 105-134 (1885); 35, 401-410 (1888); 36, 93-114 (1889).

⁸¹ Alty, T., Proc. Roy. Soc. London (A), 131, 554-564 (1931); Nature, 130, 167-168 (1932); Phil. Mag. (7), 15, 82-103 (1933).

⁸² Alty, T., and Mackay, C. A., Proc. Roy. Soc. London (A), 149, 104-116 (1935).

⁸⁸ Alty, T., and Nicoll, F. H., Can. J. Res., 4, 547-558 (1931).

of α for water decreases as the temperature increases. In that work the temperature of the surface was assumed to be that indicated by a thermojunction.

Alty and Nicoll ³³ concluded that $\alpha = 1$ for benzene (C_6H_6) at 30 °C, and R. Marcelin found $\alpha > 0.1$ for ether $[(C_6H_5)_2O]$ and for carbon disulphide (CS_2)³⁴; for nitrobenzene ($C_6H_5NO_2$), solid naphthalene ($C_{10}H_8$), and solid iodine, he found that α ranged from 0.035 to 0.25, increasing with the temperature, which was varied from 40 to 60 °C.³⁵

Alty has now extended his observations to include the case of vapors striking each its own crystalline phase, and finds that in every case examined, whether the condensed phase is liquid or crystalline, $\alpha = 1$ if the dipole moment of the substance is zero, but is small if the dipole moment is great.³⁶ Ice was not among the solids studied.

Temperature Adjustment.—It seems that, whereas, only a very small fraction of the vapor molecules striking a water surface enter it, nevertheless "all of them reach temperature equilibrium with the surface before re-evaporating" into the vapor.³⁷ These authors call the coefficient that measures the approach to such equilbrium the accommodation coefficient.

Change in Association.—In the process of changing from one phase to another there is, in general, a change in the degree of association of the molecules. The observations (p. 000) of A. W. Smith ¹¹ on the vaporization of water, and those of H. T. Barnes and W. S. Vipond ¹² on that of ice, indicate that in each case the vapor is polymeric as it leaves the denser phase, but quickly breaks down into ordinary (unassociated) water-vapor. This suggests that the change in association will usually occur in the phase into which the molecules that are being considered are entering, rather than in the one from which they come.

Evaporation.

An extensive, annotated bibliography of evaporation, chronologically arranged and covering the years 1670 to the early portion of 1909, was published by Mrs. Grace J. Livingston ³⁸; and a theoretical treatment of sublimation has been given by S. Miyamoto.³⁹

In the following, the term evaporation will be used to denote the net loss of substance from the surface of the denser phase (water or ice) in a given time, sometimes per unit surface, sometimes for a given total surface, as may appear.

Superheating.*

It is improbable that the free surface of a liquid can be heated above

- * Note is on p. 623.
- 84 Marcelin, R., J. chim. phys., 10, 680-690 (1912).
- 85 Marcelin, R., Compt. rend., 158, 1674-1676 (1914).
- 86 Alty, T., Proc. Roy. Soc. London (A), 161, 68-79 (1937) → Nature, 139, 374 (L) (1937).
- 87 Alty, T., and Mackay, C. A., Proc. Roy. Soc. London (A), 149, 104-116 (1935).
- ³⁸ Livingston, Mrs. Grace J., Monthly Weather Rev., 36, 181-186, 301-306, 375-381 (1908); 37, 68-72, 103-109, 157-160, 193-199, 248-252 (1909).
 - 89 Miyamoto, S., Trans. Faraday Soc., 29, 794-797 (1933).

the temperature at which the pressure of the vapor in equilibrium with it equals the total existing pressure on the surface, but the bulk of the liquid can be readily superheated (p. 579), and will always become superheated before boiling occurs. The increase in vaporization that accompanies the growth of the bubbles draws upon the liquid for heat, which can be supplied only if the temperature of the liquid is higher than that of the surface of the bubble. This raises the question: Does the escaping vapor have its normal temperature—the temperature at which the vapor and a plain surface of the liquid will be in equilibrium when the partial pressure of the vapor is equal to the total pressure of the existing gas phase? It is generally believed that its temperature will be normal, but certain observations by Jakob and Fritz were, at least for a time, taken as indicating that the temperature of the escaping vapor is abnormally high. interpretation seems to have rested on the assumption that the escaping vapor must have the temperature of the liquid in bulk, whereas its temperature must be that of the surface of the bubble, which is lower. F. Bošnjaković 40 has shown that the observations are entirely consistent with the vapor having its normal temperature, if each bubble may be regarded as surrounded by a layer of stagnant water 17 μ thick, through which heat passes by conduction only. H. B. Reitlinger 40a found that, when water is expanded by passing it through a suitable nozzle, it does not suddenly vaporize when its pressure has been reduced to that of the vapor saturated at the temperature of the water, but only when the pressure has been reduced still lower.

Some Factors Affecting Evaporation.

Curvature of Surface.—At a given temperature the equilibrial pressure of the vapor in contact with a concave surface of its liquid is less than that over a flat surface. If the surface is a section of a sphere of radius r, the fractional decrease in the pressure will be $\Delta p/p = 2\gamma M/\rho RTr$ approximately (p. 568), where $\gamma =$ surface tension, $\rho =$ density of the liquid. Consequently, evaporation from such a surface will be slower, and condensation upon it will be greater, than on a flat surface at the same temperature and in contact with vapor of the same density. And for a given common density of vapor there will, when possible, be a transfer of substance, by evaporation, from the flat to the concave surface when they are kept at the same temperature.

Over a convex surface the equilibrial pressure is greater than that over a flat one at the same temperature, and the several effects just mentioned are modified accordingly.

The liquid within capillary spaces is subjected to tension if the con-

^{*} Jakob, M., and Fritz, W., Techn. Mech. Thermod., 1, 173-183, 236-240 (1930); Forsch. Gebiete Ingenicuru., 2, 435-447 (1931); Jakob, M., Chem. Apparat., 19, 109-111 (1932); Schreber, K., Dinglers Polytech. J., 345, 189-191 (1930); 346, 21-27, 41-46, 61-64 (1931); Z. techn. Physik, 14, 81-85 (1933); see also, Jakob, M., Mech. Eng., 58, 643-660 (1936); Fritz, W., and Ende, W., Physik, Z., 37, 391-401 (1936); Fritz, W., and Homann, F., Idem, 37, 873-878 (1936).

 ⁴⁰ Bošnjaković, F., Techn. Mech. Thermod., 1, 358-362 (1930).
 40n Reitlinger, H. B., Compt. rend., 198, 2290-2292 (1934).

cavity of the capillary surface is directed away from the liquid (pressure. if towards), and this may modify both the coefficient of capture and the surface tension. Such effects will probably be inappreciable unless the spaces are very small. If the radius of the space were 1μ (=0.0001 cm) the tension for water would be of the order of one atmosphere. G. A. Hulett 41 has reported observations which he thought indicated that water subjected to hydrostatic tension evaporated less slowly than water not under such tension; and W. A. Patrick and N. F. Ebermann 42 have published observations which they thought indicated that the pressure of vapor in equilibrium with a very concave surface of its liquid is less than that computed by means of the formula just given, unless to y is assigned a value in excess of that found for much flatter surfaces (pp. 513 and 568). But the interpretation of each of these sets of observations is difficult. See also, p. 631, M. Polanyi, 43 and A. N. Frumkin. 48a

Blanketing Layers and Surface Films.—The rate at which vapor can actually leave the surface of a denser phase is very seriously limited by the presence of a stagnant layer of gas or of vapor that always clings to the surface, and through which the escaping vapor must pass by the slow process of diffusion. The thickness of this layer varies with the conditions. For a freely exposed surface in a wind or a stream of gas, it is estimated to be a millimeter or less.44

R. Marcelin 45 has conclude dthat, under steady conditions, the surface of a liquid has the temperature at which it will be in equilibrium with the adjacent vapor at its existing partial pressure. That is, a liquid is continually blanketed by a layer of its saturated vapor. Presumably the same is true of a solid. But it must be remembered that whenever there is an evaporative loss from the denser phase the temperature of the surface will be less than that of the bulk of the substance.

Of those who have studied the effect of thin surface films, and especially of monomolecular films, upon the rate of evaporation of a liquid may be mentioned G. Hedestrand, 46 N. K. Adam, 47 E. K. Rideal, 48 and I. and D. B. Langmuir.49 The work of Rideal and that of the Langmuirs have been discussed by T. Alty.⁵⁰ For various reasons, none of this work is suitable for an estimation of the coefficient of capture (p. 621). Hedestrand and Adam found that the presence of a monomolecular film produced very little

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41 Hulett, G. A., Z. physik. Chem., 42, 353-368 (1903).
<sup>42</sup> Patrick, W. A., and Ebermann, N. F. I. Phys'l Chem., 29, 220-228 (1925) ('f. Shereshefsky, J. L., Am. Chem. Soc., 50, 2966-2980, 2980-2985 (1928); Latham, G. H., Idem, 50, 2987-2997 (1928).
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⁴⁸ Polanyi, M., Physik. Z. Sowj., 4, 144-154 (1933).

⁴⁸a Frumkin, A. N., Idem, 4, 154-155 (1933).

⁴⁴ Jeffreys, H., Phil. Mag. (6), 35, 270-280 (1918); Giblett, M. A., Proc Roy. Soc. London (A), 99, 472-490 (1921).

⁴⁵ Marcelin, R., J. chim. phys., 10, 680-690 (1912); see also Compt. rend., 154, 587-589 (1912); 158, 1419-1421 (1914).

⁴⁶ Hedestrand, G., J. Phys'l Chem., 28, 1245-1252 (1924).

⁴⁷ Adam, N. K., Idem, 29, 610-611 (1925).

⁴⁸ Rideal, E. K., Idem, 29, 1585-1588 (1925).

⁴⁹ Langmuir, I., and D. B., Idem, 31, 1719-1731 (1927).

⁵⁰ Alty, T., Proc. Roy. Soc. London (A), 131, 554-564 (1931).

effect upon the observed rate of evaporation; that being so very much smaller (1/50000) than m_s (see Table 274), no certain conclusion can be drawn from it regarding the effect of the film upon me. At a much greater rate of evaporation (0.004m₈), Rideal found that such films produced a marked effect, in some cases reducing the rate by 50 per cent. The Langmuirs discuss these observations, interpreting them in terms of the resistance offered to the passage of the vapor through 3 distinct layers: (1) a layer of stagnant water below the film, (2) the surface film, and (3) the layer of stagnant air above the water. They suggest that the surface film itself offers little, if any, resistance to the passage of the vapor; but from its effect on the surface-tension it prevents the irregular surface streamings that would otherwise exist and that would prevent the formation of a stagnant layer of water at the surface. The effect of the film upon the rate of evaporation is, in their opinion, a secondary one, arising from the attendant formation of a stagnant layer of water, of which they estimate the thickness in one case to be 0.2 mm.

Convection.—That convection currents in the liquid may, and in most cases will, play an important role in the rate of evaporation, even when that is only a small fraction of m_s is obvious from the enormous magnitude of the temperature gradient (Table 274) that is required if the necessary heat is to be supplied solely by conduction through the liquid. See also the latter portion of the preceding paragraph.

Wind.—A wind blowing along the surface tends to sweep away the vapor, and in that way to increase the evaporation. Formulas expressing this effect will be found in Table 275, and illustrations of it in several places in this section. To the references there given, may be added M. Centnerszwer, C. Wekerówna, and Z. Majewska.⁵¹

Aspect of Surface.—Other things being the same, the evaporation from a vertical surface is nearly twice as rapid as that from a horizontal one.⁵²

Electric Charge.—J. R. Sutton ⁵⁸ has quoted various opinions and observations regarding the effect of electrically charging the vessel from which a liquid is evaporating. By actual observation, he found that the rate of evaporation was the same in all cases, whether the vessel was insulated, earthed, or charged.

Cooling by Evaporation.

The surface from which evaporation is occurring is cooler than it would otherwise be, and that on which condensation is occurring is warmer. Indeed, it is obvious from Table 274 that thermal conduction through water is totally unable to prevent a very great cooling of the surface when evaporation can proceed entirely unhindered. Even under laboratory conditions the cooling is far greater than many would expect. For example, T. Alty ⁵⁰

⁸¹ Centnerszwer, M., Wekerówna, C., and Majewska, Z., Bull. Int. Acad. Polonaise (Cracovie) (A), 1932, 369-382 (1932).

Hinchley, J. W., J. Soc. Chem. Ind., 41, 242T-246T (1922). See also Hilpert, R., Forschungsheft, 355 (1932).
 Sutton, J. R., Sci. Proc. Roy. Dublin Soc., 11, 137-178 (1907).

has found by extrapolation that when bodies of water maintained at 60, 40, and 18 °C evaporate into dry air, then the temperatures of their surfaces, as measured by means of a thermojunction, lie below those of the body of the water by 28, 22.5, and 12 °C, respectively. And H. G. Becker ⁵⁵ has reported that when water is evaporating in free air and from a vessel immersed in a bath kept at 100 °C, then the temperature of the water (not of its surface, but of the water in bulk) will be 70 °C in still air, 60 °C in a moderate draft, and only 54 °C in a strong draft; and that if the temperature of the water is to be kept at 100 °C under those air conditions, then the bath must be heated to 170, 197, and 215 °C, respectively.

Obviously, the increased cooling produced by blowing air over a water surface depends upon the initial humidity of the air, and may be increased by heating the air.⁵⁶

Table 274.—Some Data Pertaining to the Evaporation of Water See also text (p. 620+) and Table 12.

 n_s = number of molecules of saturated water-vapor that strike a flat area of 1 cm² in 1 sec; m_s is the aggregate mass of these molecules; $p_{\rm sat}$ = saturation pressure at temperature t °C; h_s = heat that must be supplied per cm²sec to compensate for the evaporation of m_s of water per cm²sec; dt/dx = temperature gradient that must exist in the water near the surface when the rate of evaporation is m_s and all the heat h_s is supplied by conduction through the water, the thermal conductivity at those gradients being assumed to be the same as under usual conditions. The values of h and of dt/dx have been computed from m_s by means of the data in Tables 272 and 130. If the unit of $p_{\rm sat}$ is 1 mm-1Ig, then n_s = $10^{21}(8.32)p_{\rm sat}/\sqrt{T}$, and m_s = $0.2476p_{\rm sat}/\sqrt{T}$ grams, T °K being the absolute temperature. (See p. 621.)

Unit of $p_{\text{sat}} = 1 \text{ mm-Hg}$; of n of $h_s = 1 \text{ kilowatt}$	$s = 10^{22}$ molecules per	(cm2sec); of m.	1 g per (cm2sec):
of $h_i = 1$ kilowatt	per cm ² ; of dt/dx	105 °C per cm =	10 °C/μ

t	PHRT	92 8	m,	h.	dt/dx
0 °C	4.58	0.23	0.069	0.17	0.31
5	6.54	0.33	0.097	0.24	0.43
10	9.21	0.46	0.136	0.33	0.58
15	12.79	0.63	0.187	0.46	0.79
20	17.54	0.85	0.254	0.62	1.06
25	23.76	1.15	0.341	0.83	1.4
30	31.82	1.52	0.453	1.10	1.8
40	55.32	2.60	0.774	1.86	3.0
50	92.5	4.3	1.28	3.0	4.8
60	149.4	6.8	2.02	4.8	7.3
80	355.1	15.7	4.68	10.8	15.7
100	760.0	32.8	9.74	22.0	

Rate of Evaporation.

As the rate of evaporation depends upon the existing conditions, which

Becker, H. G., Sci. Proc. Roy. Dublin Soc. (N. S.), 17, 241-248 (1923).
 Scott, A. W., J. Roy. Techn. Coll. Glasgow, 2, 620-629 (1932).

may vary widely, formulas of various types have been used to represent it, each assuming the existence of certain specific conditions. Some of these formulas have been assembled in Table 275; comments regarding them will be found in the accompanying notes. In general, the two phases are assumed to have a common temperature, and no attention is paid to the fact that the surface is quite significantly cooler than the rest. The surface temperature, though unknown, is definitely fixed by the conditions of the problem, and thus implicitly enters into the values found for the coefficients in the empirical equations.

The effective partial pressure (p_0) of the vapor in the layer adjacent to the liquid surface (that is, the partial pressure that must be assumed if the removal of the vapor is to be accounted for by pure diffusion) has been studied by H. Mache ⁵⁷ for the case of water contained in vertical cylinders. As the length (h) of the cylinder above the surface of the water increases, the rate of evaporation decreases, and p_0 approaches the saturation pressure corresponding to the temperature of the system (Table 276.⁵⁸

The rate of evaporation, per unit area, from large areas (lakes, etc.) is about 2/3 of that from small pans, and that of sea-water is about 5 per cent less than that of fresh water.⁵⁰

Phenomena associated with the evaporation of very small drops are considered below (p. 631). For evaporation from snow, see Table 281. To references given elsewhere, may be added M. Allen. 60

Table 275.—Formulas for the Rate of Evaporation

(Adapted from compilation by A. C. Egerton, 61 with additions.)

Quite recently, F. G. Millar ⁶² has discussed previously proposed formulas, and has derived a new one which is thought to be sounder and more generally applicable. The original paper should be consulted.

A = area of the evaporating surface, $V \equiv v_1 p_1/P$, where v_1 is the total volume of vapor (as measured at $T \circ K$ and pressure p_1) that leaves the surface in unit time, P = total pressure of the gas phase, $T \circ K =$ temperature of the system, $p_s =$ pressure of the vapor when saturated at $T \circ K$, $p_0 =$ partial pressure of the vapor in the blanket adhering to the surface, p = partial pressure of the vapor in the gas (air) to which the evaporation occurs (in that blowing over the surface, or at the upper and open end of the cylinder containing the liquid), p = distance from the surface of the liquid to the open (upper) end of the vertical cylinder containing it. p = radius of a circular cylinder or of a spherical drop, p = and p = principal semi-axes of the transverse section of an elliptical cylinder, p = velocity of gas

⁸⁷ Mache, H., Sits.-b. Akad. Wiss. Wien (Abt. IIa), 119, 1399-1423 (1910).

⁵⁸ See also, Trautz, M., and Müller, W., Ann. d. Physik (5), 22, 333-352 (1935).

⁵⁰ Egerton, A. C., Int. Crit. Tables, 5, 54 (1929).

⁶⁰ Allen, M., Proc. Nat. Acad. Sci., 10, 88-92 (1924).

⁶¹ Egerton, A. C., Int. Crit. Tables, 5, 53-55 (1929).

⁶² Millar, F. G., Can. Meteorolog. Memoirs, 1, 43-65 (1937).

Table 275 .- (Continued)

(air) parallel to the evaporating surface, D = coefficient of interdiffusion of the vapor and gas (air) corresponding to T and P, m = total mass evaporated per unit of time, m = VPM/RT where M = molecular weight of the vapor, and R = the universal gas-constant per g-mole.

 $V = C_1 D \log_e \{(P - p)/(P - p_0)\}$. If p_0 is small in comparison with P, this expression for V is essentially equal to $V = C_1 D(p_0 - p)/P$, and $m = C_1 D M(p_0 - p)/RT$. If p_0 is small and p = 0 (air perfectly dry), then $V = C_1 D p_0/P$.

Unit of a, b, r, and h = 1 cm; of A = 1 cm²; of D = 1 cm²/sec; of V = 1 cm³/sec; of w = 1 m/sec; of m as indicated

I. Theoretical formulas. Unit of m = 1 g/sec.

Gas turbulent and streaminga

Circular area, h = 0.

If w is vanishingly small, m is proportional to r; if w is finite, $m = 39.5\rho_0 (D_o w r^3)^{1/2}$ where ρ_0 = density of the vapor in the blanketing layer, and D_o = effective coefficient of diffusion.

	Gas quie	scent	
Condition ^b	h	C_1	Refo
Cylinder, Elliptical	0	$4\sqrt{ab}$	1
Cylinder, Circular	0	4r	1
Cylinder, Circular	h	$4\{(h^2+r^2)^{1/2}-h\}$	2
Cylinder, Circular	> 2r	$\pi r^2/h$	3
Cylinder, Any	> 2r	A/h	3
Concentric Spheres	•	$4\pi r_1 r_2/(r_2-r_1)$	4

II. Empirical formulas. Unit of m=1 g/hr. Circular cylinder^d: $m=1000C_2r^n$, where $C_2=5+25e^{-2h}$ and $n=2.0-0.60e^{-h}$. Flat rimless surfaces: $m=Ak_1(1+k_2rv)f$, f being a function of p, p_s , and P.

k_1	k_2	f		Ref
2.62 2.36 0.027M 0.49	0.85 0.44 2.24 2.24	$p_0(p_s - p)/P^n p_s$ $\{(p_s - p)/P\}^{1/2}$ $(p_s - p)/P$ p_s/P p_s/P	6.	Laval Hinchley Himus and Hinchley Hine Hine; for water
1.27	$\begin{array}{c} 1.12 \\ 0.063 \\ 0.293/w^{1/2} \end{array}$	$(p_{\bullet}-p)/P$	10.	FitzGerald Bigelow Grunsky
1.68	0.764	$\frac{(p_s + p_a - 2p)/P}{(p_s - p)/P}$	11. 12.	Marvin Lurie and M.

^a Derived by H. Jeffreys ^{6s} for flat circular surfaces not surrounded by an elevated rim. Of the theoretical relations that have been proposed, these probably apply the most closely to practical conditions. The value of D_{\bullet} depends upon the turbulence and convection; it should be determined experimentally in each case. That of ρ_{0} is the mass of vapor per unit volume at the top of the thin layer (1 mm or less in thickness) beyond which, on account of turbulence and convection, the concentration of the vapor decreases very slowly with the elevation. In the open, w may reach 0.4 m/sec, and D_{\bullet} reach 1000 cm²/sec; then the formula applies if $10 < r < 25\,000$ cm. Indoors, w may be 0.04 m/sec and $D_{\bullet} = 1$ cm²/sec; then the limits of validity are 1 < r < 2500 cm. (In the International Critical Tables, 5, 54 (1929) these limits are incorrectly given as areas 250 m² to 10 cm² and 25 m² to 1 cm².) The corresponding expression $m = C_{0}\rho_{0}\sqrt{D_{\bullet}w^{3}}$, in which C_{0} is a form-factor and l is a linear dimension defining the size of the surface, applies to flat surfaces of any shape, provided l Jeffreys, H., Phil. Mag. (6), 35, 270-280 (1918).

Table 275.—(Continued)

that the dimensionless quantity wl/D_e lies between 4 and 10000, the unit of w here being 1 cm/sec

^b Condition of evaporation, whether from the bottom of a vertical cylinder of height h above the surface of the liquid or from a sphere.

References and remarks. References cover both theory and observation.

1. Mache, H., or v. Pallich, J., Sitzb. Ak. Wiss. Wien (Abt. IIa), 106, 384-410 (1897); Stefan, J., Ann. d. Physik (Wied.), 17, 550-560 (1882), 41, 725-747 (1890), Sitzb. Ak. Wiss. Wien (Abt. IIa), 65, 323-363 (1872); Renner, O., Ber. deuts, bot. Ges., 29, 125-132 (1911); Winklemann, A., Ann. d. Physik (Wied.), 35, 401-410 (1888). When p = 0, po = past, and past/P is small, this expression reduces to V = 4rDpsat/P, which was given by Dalton, J., Mem. Manchester Lit. Phil.

2. Brown, H. T., and Escombe, F., Phil. Trans. (B), 193, 223-291 (1900); Laval, E., Jour. de Phys. (2), 1, 560-561 (1882) & Mém. Soc. Sci. phys. et nat. Bordeaux (2), 5, 107+ (1882); Thomas, N., and Ferguson, A., Phil. Mag. (6), 34, 308-321 (1917); Vaillant, P., Compt. rend., 150, 689-691, 1048-1051 (1910); Jour. de Phys. (5), 1, 877-891 (1911); Winklemann, A., Ann. d. Physik (Wied.), 35, 401-410 (1888).

- a. Physik (Wied.), 35, 401-410 (1888).

 3. LeBlanc, M., and Wuppermann, G., Z. physik. Chem., 91, 143-154 (1916); Marcelin, R., 45 and Compt. rend., 158, 1674-1676 (1914); Vaillant, P., Compt. rend., 150, 689-691, 1048-1051 (1910); Winklemann, A., 64

 4. Stefan, I., Sitab. Ak. Wiss. Wien (Abt. IIa), 65, 323-363 (1872), Ann. d. Physik (Wied.), 17, 550-560 (1882), 41, 725-747 (1890); Langmuir, I., Phys. Rev. (2), 12, 368-370 (1918); Houghton, H. G., Physics, 4, 419-424 (1933); Fuchs, N., Physik. Z. Sowi, 6, 224-243 (1934). Here r_1 = radius of the evaporating sphere, and r_2 = radius of a concentric spherical shell at which the partial pressure of the vapor is continuously kept equal to p. The transfer of vapor is by diffusion only, the gas being completely quiescent When $r_2 \gg r_1$, $C_1 = 4\pi r_1$, approximately, and if r_1 denotes the time, then $-d(r_1^2)/dr = 2DM(\beta_0 p)/RT$, a quantity independent of r_1 . The observed rates of evaporation of single spheres in large volumes of gas, though of low order of precision, agree with these relations; see Sresnevski, B., Zhurnal Russ. fiz.-khin. obshchestvo. 14, 420-469, 483-509 (1882), 15, 1-10 (1883); Morse, H. W., Proc. Am. Acad. Arts Sci., 45, 363-367 (1910); Gudris, N., and Kulikowa, L., Z. Physik, 25, 121-132 (1924), J. Russ. Phys. Chem. Soc. (Phys.), 56, 167-175 (1924); Whytlaw-Gray, R., and Whitaker, H., Proc. Leeds Phil. Lit. Soc., 1, 97-103 (1926); Topley, B., and Whytlaw-Gray, R., and Whitaker, H., Proc. Leeds Phil. Lit. Soc., 1, 97-103 (1926); Topley, B., and Whytlaw-Gray, R., Phil. Mag. (7), 4, 873-888 (1927); Houghton, H. G., Physics, 4, 419-424 (1933). See also p. 631.

 5. Laval, Jour. de Phys. (2), 1, 560-561 (1882) \leftarrow Mém. Soc. Sci. Phys. et Nat. Bordeaux (2),
- 5. Laval, Jour. de Phys. (2), 1, 560-561 (1882) \leftarrow Mém. Soc. Sci. Phys. et Nat. Bordeaux (2), 5, 107+ (1882), has stated that n varies with the gas, but not with the temperature; $p_0 =$ pressure of the vapor in contact with the surface.
- 6. Hinchley, J. W., J. Soc. Chem. Ind., 41, 242T-246T (1922); Himus, G. W., and Hinchley, J. W., Idem, 43, 840-845 (1924). The purpose of this work was to obtain data and formulas that would be of value to the chemical engineer. Hinchley stated that his formula applies if w > 0, and that it is not in error by more than 10 per cent if t > 60°C; whereas H. G. Becker so has stated that when w = 0 and t > 90°C then the values defined by this formula are too low. See also, Hill, L., and Hargood-Ash, D., Proc. Roy. Soc. London (B), 90, 438-447 (1919).
- 7. Hine, T. B., Phys. Rev. (2), 24, 79-91 (1924). For a circular surface 30 cm in radius. Water was not considered in setting up the formula containing M; the formula here given for water was not given by Hine, but is obtained from his formula by setting M = 18.0154. See also de Heen, P., Bull. Sci. Acad. Roy. Belg. (3), 21, 11-24 (1891)
- 8. FitzGerald, D., Trans. Am. Soc. Civ. Ena., 15, 581-646 (1886). This and the next two expressions refer to evaporation from large outdoor areas of water, ice, or snow. The value of ps for dry ice and snow is not the same as for water at the same temperature. For observed evaporation from snow, see Table 281. See also Giblett, M. A., Proc. Roy. Soc. London (A), 99, 472-490 (1921).
- 9. Bigelow, F. H., Monthly Weather Rev., 36, 24-39 (1908); it applies to the same conditions as the preceding reference (8). Marvin, C. F., Idem, 37, 57-61 (1909) regarded this value for k2 as merely a first approximation.
- 10. Grunsky, C. E., Monthly Weather Rev., 60, 2-6 (1932). (Discussion by C. F. Marvin on p. 6); it applies to the conditions stated in reference 8. His complete formula is E = E'(1 + E') $0.293\sqrt{w}$) (1 + 0.108H), unit of w=1 m/sec, of H=1 km, H= altitude above sea-level; it is intended for general use in computing the evaporation from lakes, reservoirs, etc. (see Table 278.) Note: As $k_2 w$ is 0.293 \sqrt{w} , $k_2 = 0.293/w^{0.5}$, as given in the table.
- 11. Marvin, C. F., Monthly Weather Rev., 37, 57-61 (1909). The pa in this formula is the partial pressure of the water-vapor in the air when that is saturated at its existing temperature; p/p_a is the relative humidity.
- 12. Lurie, M., and Mikhailoff, N., Ind. Eng. Chem., 28, 345-349 (1936) observed the rate of evaporation from a surface that was flush with the floor of a rectangular wind-tunnel. The total pressure (P) was always that of the atmosphere, and its value was merged with their constant C. The compiler has changed the unit of area and has taken the P out of the C, to make their expressions. sion conform to the others in this table
- In general, C_2 and n depend upon h, upon the temperature, pressure, and humidity of the air in which the cylinder is immersed, and upon the wind velocity. The values here given apply to indoor conditions (w = 0) with t = 15 to 20 °C, P = 749 to 787 mm-Hg, relative humidity = 56 to 74 per cent. [Thomas, N., and Ferguson, A., Phil. Mag. (6), 34, 308-321 (1917)].

Table 276.—Effective Partial Pressure of Blanketing Vapor 57

Water is contained in a vertical tube 2.67 mm in internal diameter, the top of the tube being h mm above the bottom of the water meniscus; the rate of evaporation into a dry gas is observed, and p_0 is the value that must be assigned to the partial pressure of the vapor at the surface of the liquid if the removal of vapor is to be accounted for by pure diffusion up the tube; t °C is the temperature of the system, assumed uniform; p_{sat} is the pressure of water-vapor that is saturated with reference to a flat water surface at temperature t; t_0 °C is the temperature at which the vapor-pressure is p_0 .

See also: A. Winklemann 64, 64a and P. Vaillant. 65

	Unit of po	and past = 1 n	nm-Hg; of h	= 1 mm. T	emp. $-t$ °C	
$\begin{array}{c} Gas \rightarrow \\ t \rightarrow \\ p_{BB} t \rightarrow \end{array}$	92.4 575.6	65.5 191.8	Air 92 4 575.6	92.4 575.6	65.5 191.8	Air 92.4 575.6
30	48.6	$-p_{\text{sat}}-p_0-$ 6.2	29.2	2.3	1.0	1.4
60 90	26.9 18.6	3.2 2.1	15.5 10.5	1.3 0.9	0.4 0.25	0.7 0.5
120 150	14.2 11.5	1.6 1.3	8 6.4	0.6 0.5	0.2 0.15	0.4 0.3

Table 277.—Various Observed Rates of Evaporation of Water

The following data are in addition to those which have been summarized by their observers in the formulas given in Table 275; w = wind velocity; m, $m_0 = \text{rate}$ of evaporation with and without wind, respectively; c = thickness of water layer removed by evaporation; c = thickness an arbitrary constant.

Unit of w = 1 m/sec; of m and $m_0 = 1$ mg/cm²hr = 0.24 mm/day; of e = 1 mm/day. Temp. = t °C

	— Becker ^a		— ← Hedestrand ^a −			_	Sutton			
70	2.54	5.0	8	crv m			South Africa			
t	n1/s	110	_	1	10.			y table 1		
50	2.8	3.8		1.5		.15		erage fo		
80	2.0	2.5		3.0	17.			.9 cm/yı		
100	1.7	2.2	2	4.5	18.	.21	= 4	1.16 mm,	/day	
Evapo	Evaporation from the oceans (Wüst ^a).									
Zon	o M		0 70	60	50	40	30	20	10	
2.011	E 14	17	0 60	50	40	30	20	10	0	
Oce	an					c —				
Atla	ıntic	0.	2 0.3	1.0	1.8	2.5	3.2	3.4	2.5	
A11	Oceans	0.	2 0.3	1.0	1.8	2.5	3.0	3.1	2.6	
Zon	. C		0 10	20	30	40	50	60	World	
Zon	6.3	1	0 20	30	40	50	60	70	Mean	
Oce	an					- c				
Atla	antic	3.	3 3.2	2.9	2.3	1.5	0.6	0.2	2.18	
A11	Oceans	2.	9 3.1	2.9	2.3	1.5	0.6	0.2	2.24	

⁶ References:

Becker, H. G.; 55 Hedestrand, G. 46 t = 20 °C; Sutton, J. R.; 55 Wüst, G., Metcor Z., 38, 188-190 (1921) (review) $\leftarrow Ver\"{o}ffentl$. Inst. Meereskunde, Berlin (N. F.) geogr.-naturw. Reihe, Heft 6; 1920, 95 S; see also Kleinschmidt, E., Meteor. Z., 38, 205-208 (1921).

⁶⁴ Winklemann. A., Ann. d. Physik (Wied.), 22, 1-31 (1884); 23, 203-227 (1884); 26, 105-134 (1885); 36, 93-114 (1889).
64 Winklemann, A., Ann. d. Physik (Wied.), 35, 401-410 (1888).
65 Vaillant, P., Jour. de Phys. (5), 1, 877-891 (1911).

Table 278.—Evaporation from Large Outdoor Areas of Water 66

 $E = E_1(1 + 0.04 \sqrt{w}) [1 + 3.3(10^{\circ}) H]$, unit of w = 1 mile/day, of H = 1 ft. $= E_1(1 + 0.293 \sqrt{w_1}) [1 + 10.8(10^{-5}) H_1]$, unit of $w_1 = 1$ m/sec, of $H_1 = 1$ m. H = altitude above sea-level; t = mean monthly temperature; w = mean monthly wind velocity; E = annual mean of the evaporation.

The value of E_1 is that given below for E when w = 0.

Unit of E=0.001 inch/day = 0.0254 mm/day; of w=1 mile/day = 0.0186 m/sec. Temp. $= t_F$ °F = t_C °C

	$zv\rightarrow$	0	100	200	300	400	500
tr	tc			E (F	H=0).		
20	-6.7	6.5	9.1	10.2	11.0	11.7	12.5
25	-3.9	9.0	12.6	14.1	15.2	16.2	17.0
30	-1.1	11.5	16.1	17.9	19.4	20.7	21.7
35	+1.7	15.0	21	23.5	25.4	20.7	28.4
40	4.4	20.0	28	31.3	33.8	36.0	37.8
45	7.2	26.5	37	41.5	44.8	47.7	50.1
50	10.0	36.0	50	56.5	60.8	64.8	68.0
55	12.8	50.0	70	78.3	84.5	90.0	94.5
60	15.6	70.5	99	110	119	127	132
65	18 3	97	136	152	164	175	184
70	21.1	127	178	199	215	229	240
75	23.9	160	224	250	271	288	302
80	26.7	196	274	307	332	353	371
85	29.5	232	325	364	392	418	438
90	32.2	270	378	423	457	486	510

[&]quot;So printed, but probably should be 27.0.

Small Drops.

See especially N. Fuchs.⁶⁷ A drop that is several microns in radius evaporates in still air in accordance with the formula given in Table 275; if r_2 is very great, then $d(r_1^2)/d\tau = -2DM(p_0-p)/\rho RT$, a quantity independent of the radius, r_1 , τ is the time, p_0 is the vapor-pressure at the surface of the drop, and ρ is the density of the liquid. See also preceding text (pp. 623 and 629). That is, the area of the surface of the drop decreases linearly with the time, if p_0 remains constant; p_0 is controllable and is assumed to be constant. But as the radius becomes smaller, of the order of 1μ (0.001 mm) or less, evaporation becomes slower; drops that in accordance with the preceding formula should vanish in a few seconds, may last for hours.⁶⁸

Gudris and Kulikowa, using drops 1μ to 0.1μ in radius, and determining the vapor-pressure at which the size of the drop remained constant, found that the saturation pressure with reference to them is that determined from the radius and the surface-tension in accordance with Kelvin's (W. Thomson's) formula (p. 568). They regarded the reduction in the rate of evaporation as the radius became smaller as an age effect, and attributed it to an absorption of the surrounding gas; it was inappreciable

⁶⁶ Grunsky, C. E., Monthly Weather Rev., 60, 2-6 (1932).

en Fuchs, N., Physik. Z. Sowj., 6, 224-243 (1934).

⁶⁶ Gudris, N., and Kulikowa, L., Z Physik, 25, 121-132 (1924); Chem. Abs., 19, 3186 (1925) ← J. Russ. Phys. Chem. Soc. (Phys.), 56, 167-175 (1924); Fuchs, N. ⁶⁷

in H₂. For other gases they published curves from which have been read the following values connecting the radius (r) of the drop with the time τ : Unit of $\tau = 1$ min: of $r = 0.01\mu = 10^{-6}$ cm

						•			
Air —			- 30 Air	+ 70 H ₂		CO ₂			
τ	*	τ	*	τ	*	τ	*	τ	7
5	94	7	99	30	84	5	89	20	86
22	89	15	94	37.5	79	10	87.5	45	86
37.5	84	22.5	89	45	74	15	86.5		

On the other hand, D. J. Woodland and E. Mack, Jr., 69 have reported observations that indicate that $-dm/d\tau = C(r + \delta)$, instead of the Cr required by the linear relation between the area and τ . They think that these observations indicate that the effective removal of the vapor occurs from a surrounding shell of saturated vapor, and not from the surface of the drop itself, the thickness of the shell being δ. Their observations lead to the values $\delta = 0.52 \,\mu$ for *n*-dibutyl phthalate, and $\delta = 1.1 \,\mu$ for *n*-dibutyl tartrate.

C. Barus 70 has stated that, in dust-free air saturated with water-vapor and left undisturbed, the dissipation of very small fog particles "by evaporation is enormously more important than by subsidence"; in his case the two were about equal when the diameter (d) of the particle was 3μ . "Fog particles precipitated on solutional nuclei (phosphorus) evaporate" to water nuclei which persist without other loss than by subsidence. Those precipitated on nuclei of water-vapor evaporate almost without residue, the persisting nuclei being only 0.4 per cent when $d = 1.6 \mu$, and 3.6 per cent when $d = 32 \mu$. "These fog particles vanish into the wet air from which they were precipitated and the experiment may be repeated indefinitely. Relatively more water nuclei persist as the fog particles evaporated are larger."

The evaporation of drops in a stream of air of velocity w m/sec has been studied by T. Namekawa and T. Takahashi 71 who found that $dr/d\tau =$ $-2.53 [(p_0 - p)/r] \cdot [1 + 2.1w^{1/2}] \cdot 10^{-7}$ if r < 1 mm and w < 2 m/sec. (See also Y. Takahasi ⁷² and E. G. Zak. ⁷²)

Condensation.

For those condensation data and phenomena in which condensation may be considered as merely negative vaporization, reference should be made to the preceding pages devoted to vaporization, one exception being Table 281 treating of condensation upon snow.

When water-vapor condenses on a surface that is chilled to -110 °C or lower, the deposited ice is vitreous.⁷⁸ Condensation on extended surfaces at higher temperatures, depending, as it does, upon both the nature

⁶⁹ Woodland, D. J., and Mack, E., Jr., J. Am. Chem. Soc., 55, 3149-3161 (1933).

⁷⁰ Barus, C., Am. J. Sci. (4), 25, 409-412 (1908).

n Namekawa, T., and Takahashi, T., Mem. Coll. Sci. Kyoto (A), 20, 139-146 (1937).

⁷² Takahasi, Y., Sci. Abs. (A), 40, 265 (1937) ← Geophys. Mag., Tokyo, 10, 321-330 (1936); Zak, E. G., Chem. Abs., 31, 3360 (1937) ← Zhur. Geofix., 6, 452-465; 466-473 (1936).

⁷⁸ Burton, E. F., and Oliver, W. F., Proc. Roy. Soc. London (A), 153, 166-172 (1936) → Nature, 135, 505-506 (L) (1935).

of the surface and local peculiarities thereof, is not considered in this compilation.

Supersaturation.—It is probable that water-vapor can never become supersaturated (supercooled) in the immediate presence of water or of ice, but when no condensed phase is present it can be considerably supercooled if it contains no dust or other nucleus on which condensation can begin. See the following section: Nuclear Condensation. Supercooling is always understood to be that with reference to vapor in equilibrium with a flat surface of water.

C. F. Powell ⁷⁴ has remarked that Callendar and Nicholson ^{74a} pointed out that the steam in the cylinder of a steam engine might be supersaturated, and that supersaturated steam might exist in a steam turbine, as has since been pointed out by others. "According to Callendar, the steam passes through the turbine so quickly that thermal equilibrium cannot be maintained. It becomes supersaturated, and no appreciable condensation takes place until the cloud-limit is reached, when nuclei are produced in enormous numbers." (Cloud-limit = supersaturation at which condensation on uncharged nuclei begins. See Table 279.) He comments on erroneous values assumed by Callendar and by H. M. Martin for the supersaturation at the cloud-limit, and from his own observations computes the data in Table 280.

When steam is expanded by passage through a simple convergent-divergent nozzle, the steam becomes supersaturated; condensation does not occur until the steam has reached the condition approximately represented by th 3.5 per cent moisture line on the Mollier enthalpy-entropy diagram, and then drops 6.2A in radius are formed.⁷⁵ For the corresponding phenomenon for water, see preceding text (p. 623).

Nuclear Condensation.—In the study of nuclear condensation, the customary procedure is to cool a mixture of gas and vapor by an adiabatic expansion, the amount of expansion that just suffices to produce condensation being determined by trial. The expansion (E) is defined as the ratio of the expanded volume (v_2) to the volume (v_1) before expansion $(E = v_2/v_1)$. If $\gamma = c_p/c_v = \text{ratio}$ of the principal specific heats of the gas-vapor mixture, and if T_1 °K is the absolute temperature of the mixture before expansion, then the temperature (T_2) °K after expansion will be given by the relation $T_1/T_2 = (v_2/v_1)^{\gamma-1}$. The conditions are usually such that the vapor in the mixture of volume v_1 and temperature v_2 is saturated with reference to a flat surface of its liquid. The corresponding vapor pressure v_2 is pressure v_3 of the vapor after expansion is given by the relation $v_2/v_1 = (v_1/v_2)^{\gamma}$; and vapor pressure $v_2/v_1 = (v_1/v_2)^{\gamma}$ corresponding to saturation at $v_2/v_1 = (v_1/v_2)^{\gamma}$; and vapor pressure $v_2/v_1 = (v_1/v_2)^{\gamma}$ corresponding to saturation at $v_2/v_1 = (v_1/v_2)^{\gamma}$.

⁷⁴ Powell, C. F., *Proc. Roy. Soc. London (A)*, 119, 553-577 (1928).

⁷⁴ Callendar, H. L., and Nicolson, J. T., *Min. Proc. Inst. Civ. Eng. (London)*, 131, 147-206-268 (1897).

W Yellott, J. I., Trans. Am. Soc. Mech. Eng., 56, 411-427-430 (FSP-56-7) (1934); Yellott, J. I., and Holland, C. K., Idem, 59, 171-183 (FSP-59-5) (1937); Jakob, M., Z. techn. Physik, 16, 83-86 (1935).

found from tables. The ratio $S = p_e/p_2$ is called the supersaturation produced by the expansion. The value of the expansion (E) depends only on the volumes v_1 and v_2 , but that of S depends also upon the nature of the inert gas and of the vapor.

As E is gradually increased, an initial and four other stages of condensation can be distinguished. Starting with air taken directly from the atmosphere and saturated, one obtains a dense cloud of drops when E is

Table 279.—Condensation of Water-vapor on Nuclei

E= expansion, S= supersaturation, each at the beginning of condensation on nuclei of the nature indicated by the subscript (-= negative ions, += positive ions, 0= uncharged nuclei). For exact definition of E and of S, see text. The values of S are unaffected by the nature of the admixed uncondensible gas; those for E refer to vapor mixed with air; t_1 and t_2 °C = temperature before and after expansion, respectively.

T. Adapte	d from L. F	B. Loeb (IC	$T)^{8}$; $t_1 =$	18 °C.		
E_	S_	E_{\star}	S_{+}	E_{0}	S_0	References ^a
1.25 1.29	4.15	1.31	5.8	1.38 1.42	7.9	Wilson Donnan
1.265 1.270 1.251		1.314 1.32		1.366 1.31	6	Przibram Andrén Laby

II. C. F. Powell (1928). 70 Values corrected for evaporation from walls, see text. E_{-} S_{-} t_1 7 -26.48.95 1.375 7 7.80 3.98 18 1.245 -6.51.370 -16.418 35° 1.314 5.07 +10.03.44 35 + 3.2 1.235 +19.150 3.74 50 1.226 +24.72.96 1.2862.52 1.252 +47.02.87 1.218 +50.5

III. Volmer and Flood.^a Values averaged by the observer over ranges of 2 °C or less.

t_1	E_{-}	t_2	S_0	t_1	E_{-}	t_2	S_0
29.2 18.	1.266 1.276	1.7 -9.4	4.18 4.85	14.7	1.288	-12.2	4.98

a References:

Andrén, L., Ann. d. Physik (4), 52, 1-71 (1917); Anderson, E. X., and Froemke, J. A., Z. physik. Chem. (A), 142, 321-350 (1929); Donnan, F. G., Phil. Mag. (6), 3, 305-310 (1902); Laby, T. H., Phil. Trans. (A), 208, 445-474 (1908); Powell, C. F.; Przibram, K., Jahrb. Radioak., 8, 285-308 (1911) Bibliography of 135 titles; Volmer, M., and Flood, H., Z. physik. Chem. (A), 170, 273-285 (1934); Wilson, C. T. R. 788

^b E. X. Anderson and J. A. Froemke (*loc. cit.*) found $E_{-}=1.201$ and $S_{-}=3.0$ at $t_{1}=25$ °C.

only slightly greater than unity. Allowing these to subside, and repeating the process, and so continuing, one presently reaches a stage at which such small expansions produce no condensation. This terminates the initial stage.

Continuing the process with gradually increasing values of E, no condensation (except on the walls) occurs until E = 1.25, when a few drops of rain form in the interior of the gas. This is stage 1. A further increase

in E merely increases the number of drops—the heaviness of the rain—until E=1.38 (stage 2). Beyond E=1.38 a persistent cloud of small drops is formed (stage 3); the number of drops increases rapidly as E increases beyond 1.38, each drop becoming correspondingly smaller, and presently diffraction colors border the image of a source of light seen through the cloud, and change as the size of the drops decreases (stage 4).

In the initial stage—generally not counted, but regarded as a cleansing process—condensation occurs on dust and other, presumably large, nuclei that are not being continually replaced. Repeated condensations sweep these out of the gas. At E=1.25 condensation begins to occur on negative ions; at higher values, on positive ions; and at E=1.38 on uncharged nuclei which "have been identified with the associated molecules present in water-vapour."

Powell ⁷⁶ found that in general the expansion actually realized is not simply adiabatic, but is attended by evaporation from the walls of the vessel, which causes the density of the expanded vapor to exceed that corresponding to simple expansion. He corrected his observations for this effect; the values so corrected are given in Table 280. He has concluded that ordinary room temperatures are to be preferred to others as initial temperatures in the investigation of atomic phenomena by means of the cloud method.

Table 280.—State of Water-vapor at the Cloud-limit 76

Values above 47 °C were obtained by extrapolation based on $S_0=1$ at the critical point. Callendar's equations for steam were used. H= heat content, P= vapor pressure, $S_0=$ supersaturation $=P_0/P_{\rm sat},~V=$ specific volume, $\phi=$ entropy, the subscripts $_0$ and $_{\rm sat}$ indicate, respectively, that the value is that at which condensation on uncharged nuclei begins (the cloud-limit), and that corresponding to equilibrium with a flat surface of water. As usual, H and ϕ are measured from water at 0 °C.

Unit of $P = 1 \text{ 1b*/in}^2 = 0.0680$	atm = 68.95 kdvne/cm2; of	V=1 f	$t^3/lb = 62.429$	cm3/g;	of
H = 1 lb-cal/lb = 1 g-cal/g	; of $\phi = 1$ lb-cal/lb °K = 1	g-cal/g.	K. Temp	t °C.	

ŧ	Prat	So	Vest	V_0	P_0	H_0	ϕ_0
	°C 0.0892	5.40	3275.9	606.5	0.4811	594.02ª	1.9899
10		4.40	1693.8	584.9	0.7855	598.72°	1.9529
20		3.70	922.19	249.2	1.254	603.32°	1.9174
30		3.25	525.8	161.8	1.992	607.85°	1.8819
40	1.070	3.02	312.4	[*] 103.5	3.214	612.30°	1.8441
50	1.789	2.81	192.7	68.59	4.995	616.67ª	1.8098
60	2.887	2.35	122.9	48.59	7.250	620.98°	1.7821
75	5.586	2.22	66.20	29.82	12.31	627.21°	1.7448
90	10.161	1.99	37.81	19.00	19.99	633.19°	1.7074
100		1.86	26.79	14.40	26.96	637.02	1.6872
120		1.66	14.27	8.595	47.11	644.22	1.6450
140		1.51	8.143	5.392	78.91	650.70	1.6068
160		1.39	4.923	3.542	122.7	656.85	1.5755
180		1.29	3.127	2.408	185.6	662.24	1.5423
200	225.2	1.20	2.074	1.728	266.0	667.58	1.5205
a	For $P_0 < 20$,	the values	of H_0 fall	on the line	of 2 per	cent wetness.	

⁷⁶ Powell, C. F., Proc. Roy. Soc. London (A), 119, 553-577 (1928). See also Anderson, E. X., and Froemke, J. A., Z. physik. Chem. (A), 142, 321-350 (1929).

Optical methods for determining the size of suspended water droplets have been discussed by J. G. Wilson 77 who has concluded that the radii (r) of the drops responsible for the colors observed by C. T. R. Wilson 78 when water-vapor was condensed by expansion (E) were as follows, the unit being $0.01 \mu = 10^{-6}$ cm: Brilliant green, E 1.412, r 135; blue-green, E 1.416, r 130; brilliant blue, E 1.418, r 119; purple, E 1.420, r 105; red, E 1.426, F 84; reddish yellow, E 1.430, F 77; orange-white, E 1.436, F 60; whitish, E 1.448, F 48; greenish white, E 1.454, F E 45.

These estimates of the sizes of the drops that cause the observed diffraction effects should not be confused with those that C. T. R. Wilson 78 has made of the equivalent sizes of the nuclei upon which the condensations began.

M. Akiyama ⁷⁹ has reported that 50 per cent of the charged recoil atoms of actinium-A do not act as nuclei of condensation for water-vapor at super-saturations in the neighborhood of those at which the vapor condenses on ordinary positive ions.

P. I. Dee 80 has published a diagram that facilitates the determinations of the quantities required in the interpretation of such adiabatic expansions of air saturated with water-vapor as are here considered.

G. Stüve 81 has concluded that in natural atmospheric condensation, gaseous nuclei give only drops of water, nuclei consisting of soluble salts give drops if condensation begins at temperatures above -20 °C, but stars of snow if below -20 °C, and insoluble hygroscopic nuclei give needles of ice at all temperatures below 0 °C.

Much is yet to be learned about the natural condensation of atmospheric moisture. Fogs may occur in air that is unsaturated; they may be absent from air that is saturated with water-vapor. The nature of atmospheric nuclei of condensation has been considered by Bennett, H. Landsberg, J. H. Coste and H. L. Wright, C. Junge, and H. Köhler, to mention only those that have happened to come to my attention. Salt from the ocean is generally believed to be the most abundant of the natural nuclei, but human activities—fires, furnaces, etc.—contribute droplets of nitreous acid and probably some of sulphuric acid. It has been frequently reported that the volumes of the drops of rain, and perhaps of fog also, are simple multiples of a few primary sizes; which suggests that the larger ones are formed by the coalescence of the smaller ones. The suggests are simple of the coalescence of the smaller ones.

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77 Wilson, J. G., Proc. Cambridge Phil. Soc., 32, 493-498 (1936).

78 Wilson, C. T. R., Phil. Trans. (A), 189, 265-307 (282) (1897).

79 Akiyama, M., Compt. rend., 187, 341-342 (1928).

70 Dee, P. I., Proc. Cambridge Phil. Soc., 28, 93-98 (1932).

81 Stüve, G., Gerlands Beitr. zu Geophys. (Kōppen Bd. 1), 32, 326-335 (1931).

82 Bennett, M. G., Sci. Abstr. (A), 37, 259 (1934) ← J. Roy. Meteor. Soc., 60, 3-14 (1934).

83 Landsberg, H., Monthly Weather Rev., 62, 442-445 (1934).

84 Coste, J. II., and Wright, II. I., Phil. Mag. (7), 20, 209-234 (1935).

85 Junge, C., Gerlands Beitr. zu Geophys., 46, 108-129 (1935).

86 Köhler, II., Arkiv. Mat., Astron., Fysik, 24 A, No. 9 (1934).

87 Gold, E., Nature, 133, 102 (L) (1934); Köhler, H., Trans. Faraday Soc., 32, 1152-1161 (1936); Markl, E., Meteor. Z., 54, 173-183 (1937).

88 Loeb, L. B., Int. Crit. Tab., 6, 117 (1929).
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Table 281.—Condensation on Snow in the Open 88a

Observations at high latitudes; data are as applicable to evaporation as to condensation. C = amount of condensation in the time τ , p = partial pressure of water-vapor in the air, $p_{\text{sat}} =$ pressure of water-vapor saturated with reference to the surface of the snow. Rolf ^{88a} finds $C = a + b(p - p_{\text{sat}})\tau$; the values of a and b vary with the season (mean temperature?), and if the ground is only partly covered with snow, with the time of day; t is the temperature of the air. The observations were admittedly rough, and in some cases differed by 100 per cent from the values defined by his equation.

```
Unit of C = 1 mm of water, of \tau = 1 hr, of p and p_{sat} = 1 mm-Hg
   Winter, ground covered, t=+0.9 to -27.5 °C, a=0, b=0.0174 Spring, ground covered, t=+3.1 to +8.7 °C, a=-0.0010, b=0.0168. Summer, ground partly covered, a and b variable as follows:
    Time of day
                            1000a
                                               1000b
                                                                 Time of day
                                                                                                            1000b
                                                                                           1000a
24 \pm 3
25 \pm 2
15.7 \pm 1.1
                                         +29 \pm 1
                                                             15:15 to 17:00
                                                                                       20 \pm 3
                                              28 \pm 2
                                                             17:15 to 20:00
20:15 to 8:10
                                                                                      8 \pm 2
                            16 \pm 4
                                              29 \pm 3
                                                                                         0.6 \pm 1.2
```

Condensation on Metals.—The conclusion of O. Reynolds ⁸⁹ that "there is no limit to the rate at which pure steam will condense but the power of the surface to carry off the heat" is practically correct, but, as he pointed out, when an uncondensing gas is mixed with the steam, the surface on which the condensation occurs becomes blanketed with a layer of gas that is relatively poor in steam, and through which the steam must pass by diffusion in order to reach the surface. That greatly reduces the rate of condensation. Furthermore, condensation may occur either as isolated drops or as a continuous film of water, depending upon the surface conditions. ⁹⁰ Phenomena relating to the condensation of steam while flowing through cooled metal tubes, though of great technical importance, scarcely fall within the scope of this compilation.

97. Freezing and Melting

(See also Section 59. For external work and change in volume on freezing or melting, and for latent heat, see Section 95; for melting temperature, Section 92.)

Ice Needles.

As freezing continues after a volume of water has become completely surrounded by ice, or enclosed between rigid walls and an ice-sheet adherent to those walls, the pressure of the water increases, and may rupture the bounding sheet of ice. If the freezing is proceeding very rapidly, as when the water is much supercooled before the freezing begins, the rupture may consist of one or more small perforations. The peripheral portions of the

⁸⁸a Rolf, B., Arkiv. Mat., Astron., Fysik, 9, No. 35 (1914).

⁸⁹ Reynolds, O., Proc. Roy. Soc. London (A), 21, 275-283 (1873).

⁹⁰ Schmidt, E., Schurig, W., and Sellschop, W., Tech. Mechan. Thermod., 1, 53-63 (1930).

issuing water will freeze, thus building up a tube of ice, which may grow with surprising speed, sometimes straight but more often curved or abruptly bent, and may attain a length of several centimeters. When the growing ceases, the contents of the tube freezes, converting the whole into a needle of ice. Such growths are not at all uncommon when there has been much supercooling. Their growth is of exactly the type described by H. Erlenmeyer 90a in explaining the growth of hair-like crystals sometimes observed to form in salt crystals that are creeping over solid surfaces. It seems probable that the "long crystal" mentioned by T. Alty 90b and the unusual one reported by O. Bally 91 were formed in this way; and the same may have been true of some of those reported by J. Meyer and W. Pfaff. 92 But in the case they mention, of a needle that grew from the wall of an empty bulb connected to one containing water, it seems that the growth must have been by condensation, as they suggest. In view of the low thermal conductivity of ice and of water-vapor, and of the large amount of heat liberated when watervapor is converted to ice, it would seem that the growth of a needle by such condensation would be extremely slow. Furthermore, the large specific volume of the vapor would restrict the size of a needle, formed by condensation, to a small value, whenever the growth occurred after the water had become completely covered with ice. For example, the volume of the needle that can be produced by the complete freezing of 1 cm³ of vapor saturated with reference to water at -15 °C (sp. vol. = 622 000 cm³/g) cannot exceed that of a cylindrical needle 0.1 mm in diameter by 0.49 mm in length; since, owing to the vapor pressure of ice at -15 °C, only about 2.2 per cent of the total volume of the vapor can so freeze, there would have to be available 45 cm³ of the vapor saturated with reference to water at -15 °C if a spicule of that small size (0.1 mm by 0.49 mm) is to be actually formed in that manner.

It is not uncommon for such needles to be spoken of as monocrystals, but the compiler has yet to see experimental evidence justifying such a description.

Supercooling of Water.

[NOTE: Since this section was written, G. Tammann and A. Büchner 93 and J. Meyer and W. Pfaff 92 have studied the subject, and have interpreted their observations in terms of the ideas that they had developed in the course of their studies of other substances. The two groups differ mainly in that Meyer and Pfaff hold that the primordial nuclei upon which the ice is first formed are always solids, foreign to the molten substance, whereas Tammann and Büchner think that they may be formed from the molten substance itself. Each group seems to think that the growth of, or

⁹⁰a Erlenmeyer, H., Helv. Chim. Acta, 13, 1006-1008 (1930).

⁹⁰b Alty, T., Phil. Mag. (7), 15, 82-103 (1933).

Bally, O., Helv. Chim. Acta, 18, 475-476 (1935).
 Meyer, J., and Pfaff, W., Z. anorg. allgem. Chem., 224, 305-314 (1935).

⁹⁸ Tammann, G., and Büchner, A., Z. anorg. allgem. Chem., 222, 371-381 (1935).

upon, the nucleus is slow until a certain critical size is attained; then it becomes rapid—visible freezing occurs. If this were true, then the time that elapses between the instant that a specimen of water of given form and size is immersed in a cold bath and the instant that visible freezing begins would be a matter of prime importance; and that time, or its equivalent, is observed and reported by each. None of this necessitates a rewriting of this section.

But more recent observations by the compiler have shown that certain of the impressions conveyed by this section are entirely wrong, although in entire harmony with the reports cited. On account of that harmony it has seemed desirable to let the section stand as written, and to request the reader to bear in mind the following facts, ascertained by the compiler.

When water is protected from the atmosphere, as by sealing it in a glass bulb, it freezes spontaneously at a fixed temperature that is independent both of the rate of cooling and of the time that the bulb has been held at a low temperature. The temperature of this spontaneous freezing varies from specimen to specimen, and for any one specimen may exhibit oscillations and may drift, but in many cases it remains constant within a few tenths of a degree centigrade for months at a time. The volume of each specimen used was about 8 cm³, and it was contained in a bulb of about twice that size. It is not at all difficult to obtain specimens of this size that can be supercooled to -14 °C, and one has been repeatedly cooled to -21 °C. In no case was any precaution taken to keep the water quiescent; in fact, the supercooled water can be poured with impunity over the entire interior of the bulb. Violent splashing will cause the supercooled water to freeze, but sharp rapping of the exterior of the bulb is without effect. On the other hand, a very gentle wiping of the glass-water interface will cause freezing at a temperature well above that at which the specimen freezes spontaneously. All the observations so far obtained are consistent with the opinion that, within the range of temperature covered (0 to -21 °C), the presence of foreign solids is essential to the spontaneous freezing of the specimen; in this respect they agree with the conclusions of Meyer and Pfaff. They also show conclusively that the time required for a specimen to freeze when subjected suddenly to a given condition of chilling is not a factor of prime importance; it is the temperature, that is reached by some portion of the specimen, that determines the freezing; and that temperature is a characteristic of the specimen, varying from one to another as do the foreign solids that serve as "nuclei." 94]

Under suitable conditions, water can be cooled below its so-called freezing point (the melting point of ice) without becoming solidified. In that state it is said to be supercooled. If a bit of ice, no matter how minute, be touched to supercooled water, freezing on the surface of the ice begins at once, and proceeds rapidly until enough latent heat has been freed to raise the temperature to the so-called freezing point. It then stops unless heat is being abstracted from the mixture.

See Dorsey, N. E., J. Res. Nat. Bur. Stand., 20, 799-808 (RP1105) (1938).

Such seeding with a suitable crystal seems to be the only means by which the freezing of supercooled water can be initiated at will. And that freezing is simply a growth of ice from the crystal into the water. The various other methods that have been suggested, some of which are mentioned below, are so variable in their action and have been shown to fail so signally in certain instances that one cannot seriously regard any of them as having been shown to be more than secondarily involved in the initiation of such freezing as may have occurred when it was employed. Under some conditions it may have facilitated the initiation, but it can scarcely be regarded as having been primarily involved in the initiation itself.

There are, however, certain conditions that appear to favor the supercooling of water, to reduce the likelihood of freezing being initiated. A. Mousson 95 reported that water can be more readily supercooled when it exists (1) as small drops on surfaces that are not wetted (velvet, finely dusted surfaces, certain leaves, etc.), (2) bubbles [?] in a fog, (3) in narrow capillary tubes, (4) as a thin layer between glass plates that are clamped together (if the plates were not clamped, if the upper one was merely laid upon the under, the water between them froze), than when it exists in bulk; and when protected from mechanical shocks, than when subjected to them. He drew the general conclusion that whatever impedes a rearrangement of the particles of the water facilitates its supercooling. L. Dufour 96 reported that the placing of a layer of oil upon the surface of the water is a very uncertain means for facilitating supercooling, but that drops of water, especially when small, suspended in a liquid of the same density could be markedly supercooled, even when subjected to violent deformations; he occasionally carried it to -20 °C. S. W. Young and R. J. Cross 97 have stated that long-continued heating facilitates the supercooling of a liquid.

In his discussion of the subject, Dufour ⁹⁶ seems to have failed to distinguish between the removal of that which impedes and the imposition of that which causes. The result of an effective impediment is the same whether or no a cause otherwise efficacious is present, but in the absence of such a cause, the removal of the impediment changes nothing. Many later investigators of supercooling have likewise failed to see the distinction, and much confusion has resulted. In the following paragraphs an attempt is made to give the opinions and the points of view of the several authors cited; the reader should constantly bear in mind the distinction just drawn, and interpret the statements accordingly.

H. T. Barnes 98 has stated that agitation, with the presence of dust or suspended matter and, particularly, of dissolved air, makes supercooling almost impossible. With the avoidance of those conditions he has cooled

⁹⁵ Mousson, A., Ann. d. Physik (Pogg.), 105, 161-174 (1858).

⁹⁰ Dufour, L., Arch. des sci. phys. et nat. (N.P.), 10, 346-371 (1861) → Ann. d. Physik (Poog.), 114, 530-554 (1861); letter appended to Arch. des sci. phys. et nat. (N.P.), 11, 22-30 (1861).

on Young, S. W., and Cross, R. J., J. Am. Chem. Soc., 33, 1375-1388 (1911).

⁹⁸ Barnes, H. T., "Ice Formation," pp. 95-97, 1906.

water in open flasks to -6 °C. He writes: "Curiously enough, once the freezing point was passed it was certain that several degrees below that point would be reached without ice forming, and at -3° or -4° quite violent agitation was required for solidification to take place It seems harder to pass the freezing point without ice forming than to continue the cooling beyond this temperature. The degree of instability reaches such a critical state, however, beyond five or six degrees, that extraordinary precautions have to be taken for further cooling."

This relative stability of the slightly supercooled water as compared with the great instability of that cooled to $-6\,^{\circ}\text{C}$ or lower led to the suggestion that a metastable condition, in which crystallization can be initiated only by seeding with ice, existed for a few degrees below zero, and that at lower temperatures a truly labile condition existed. But S. W. Young, ⁹⁹ S. W. Young and R. J. Cross, ¹⁰⁰ and S. W. Young and W. J. von Sicklen ^{100a} seem to have shown that no such distinction exists. The difference is merely one of degree; the whole supercooled "field is labile and crystallization may be brought about in any portion of it by the production of sufficient mechanical shock." Young and von Sicklen found freezing to attend mechanical shock when the temperature of the water was as high as $-0.02\,^{\circ}\text{C}$.

Observations reported in the three papers just mentioned, and especially those in the first, indicate that the frequency with which the freezing of supercooled water accompanied the friction of solid on solid within the water, varied with the nature of the solid.

Contrary to the observations of H. T. Barnes 98 are those of R. Pictet (mentioned by Oltramare 101), who found that supercooled water in a stoppered flask can be violently agitated without its freezing, even though the temperature be -19 °C. The flask was half full. The present compiler has observed the same for a few cubic centimeters of supercooled water at -14 °C in a sealed bulb of about twice the volume of the water.

Observations on the supercooling of drops of water suspended in a liquid of the same density, and of water in capillary tubes have been reported by T. Borovik-Romanova, 102 who gave the following values for the temperature (t) at which freezing began in capillary tubes (presumably of glass) of diameter d, and for the ranges over which the observed values were spread:

At a much earlier date, H. C. Sorby 103 had made similar observations, carrying the supercooling in tubes to -16 °C, and observing, partly in

⁹⁹ Young, S. W., J. Am. Chem. Soc., 33, 148-162 (1911).

¹⁰⁰ Young, S. W., and Cross, R. J., Idem, 33, 1375-1388 (1911).

¹⁰⁰a Young, S. W., and von Sicklen, W. J., Idem, 35, 1067-1078 (1913).

on Oltramare, G., Arch. sci. phys. et. nat. Genève (3), 1, 487-501 (1879).

 ¹⁰⁸ Borovik-Romanova, T., Chem. Abs., 19, 3186 (1925) ← J. Russ. Phys. Chem. Soc. (Phys. Part), 56, 14-22 (1924) (Russian).
 108 Sorby, H. C., Phil. Mag. (4), 18, 105-108 (1859).

company with Tyndall, that even at temperatures near $-20\,^{\circ}\text{C}$ the liquid* occurring in natural cavities in quartz does not freeze, the diameter of the cavity being about 0.25 mm (0.01 in.). He observed further, that even in capillary tubes water will freeze at a temperature very near 0 °C if the water is at any point in contact with ice, and that it melts at 0 °C. And C. Despretz, 104 using water thermometers with bulbs several lignes in diameter (1 ligne = 2.25+ mm), had followed the dilatation of water (unfrozen) to $-20\,^{\circ}\text{C}$. He stated that Blagden (no citation) had cooled water to $-6\,^{\circ}\text{C}$, and Gay-Lussac (no citation) to $-12\,^{\circ}\text{C}$. G. Oltramare 101 has stated, without citation, that both Pictet and Dufour had cooled water to $-40\,^{\circ}\text{C}$. This is the greatest supercooling that the compiler has found mentioned; he has not succeeded in finding a paper by either Pictet or Dufour reporting this value.

Of the more recent investigations of supercooling, may be mentioned the following: II. A. Miers and Miss F. Isaac, 106 using water sealed in tubes "which were vigorously and continuously shaken by hand in a bath of brine" cooled at the rate of 2 °C per hour, found freezing to occur between -1.6 and -2 °C, averaging -1.9 °C, at which temperature, they concluded, "pure water freezes spontaneously, *i.c.*, in the absence of ice particles." They remark that the index of refraction is a maximum at about the same temperature.† When the tubes contained loose bits of solids (glass, garnet, lead) rubbing with friction on the walls as the tubes were shaken, freezing might occur as high as -0.4 °C. W. H. Martin 107 has reported that water that had been repeatedly redistilled *in vacuo* and without ebullition could be cooled in a 2-mm tube to -26 °C, whereas ordinary distilled water froze at -11 °C under the same cooling conditions. G. V. Lange 108 has cooled water in 0.1 mm capillaries to -18 °C, the cooling having taken 10 hours.

That very small water droplets suspended in the air may be cooled to very low temperatures without freezing, is indicated by the well-known fact "that the most brilliant coronas—those of multiple rings and large diameter—usually are formed by very high clouds whose temperature often must be far below freezing." ¹⁰⁹

L. Hawkes 110 has stated that the deposit on the cooling pipes in a room maintained at -17 to -22 °C "was found to be a mixture of water drops

^{*}Sir Humphry Davy 105 had previously found that every such clear, colorless, liquid inclusion which he had examined consisted of nearly pure water, and with few exceptions, was under less than atmospheric pressure.

[†] The maximum value of the index probably lies much nearer 0 °C, see p. 280+.

¹⁰⁴ Despretz, C., Ann. chim. phys. (2), 70, 5-81 (1839) → Compt. rend., 4, 124-130 (1837) → Ann. d. Physik (Pogg.), 41, 58-71 (1837).

¹⁰⁶ Days, Sir Humphry, Phil. Trans. 112, 367-376 (1822) = Ann. chim. phys., 21, 132-143 (1822) = Annals Philos. (N. S.), 5, 43-49 (1823).

¹⁰⁶ Miers, H. A., and Isaac, Miss F., Report Brit. Assoc. Adv. Sci., 1906, 522, (1906).

¹⁰⁷ Martin, W. H., Trans. Roy. Soc. Canada, III (3), 7, 219-220 (1913).

¹⁰⁸ Lange, G. V., Jour. de phys. (7), 1, 406D (1930) ← Bull. de l'Inst. Agronom, Kharkow, 8-9, 107-108 (1929).

¹⁰⁰ Humphreys, W. J., "Physics of the Air," 2nd ed., p. 534, 1929.

¹¹⁰ Hawkes, L., Nature, 124, 225-226 (1929).

and ice—this at -22 °C." The statement seems to imply that the water drops were at -22 °C, although in contact with ice; such surely was not the case. The presence of drops indicates continuing condensation, and had the temperature of the drops been directly observed there is no doubt that it would have been found to lie above 0 °C.

Both this and his preceding note 111 are entitled "Super-cooled Water," and refer to the apparently vitreous solid that Beilby obtained when a small drop of water was rapidly chilled to a temperature some 15 degrees below 0 °C (see p. 396). That is not the state of supercooling with which we are here concerned. We are now concerned solely with water in its fluid state.

The freezing of water in such capillary systems as soils, sand, and silica has been discussed by E. A. Fisher. 112

Superheating of Ice.

Although water can be supercooled, there is no evidence that ice can be superheated—heated above its melting point (pp. 405 and 604). Nevertheless, the temperature of a well-stirred intimate mixture of ice and water will differ slightly from 0 °C if heat is either withdrawn or supplied very rapidly. H. T. Barnes 113 states that "the ice itself shares in the temperature elevation or depression" and explains the departure from 0 °C by the inability of the ice "to freeze or melt rapidly enough to keep up the heat The velocity of crystallization and of melting is finite, and is the determining factor in the temperature of the two phases when coexisting." But that the temperature of the ice itself should rise above 0 °C under such circumstances seems most improbable, and is, indeed, contrary to the general experience which Barnes 113, p. 90 expresses thus: "It seems to be impossible to superheat a solid with respect to a liquid." It is more likely that the ice does not rise above zero, and that it is protected from the action of the surrounding water at higher temperature by a thin, closely adherent blanket of colder water. In the reverse case, a corresponding blanket of warmer water, heated by the latent heat freed as the water freezes, will protect the ice from excessive chilling, and may keep it at zero even though the temperature of the bulk of the water is slightly lower. It seems probable that in the case of ordinary ice (ice-I) the actual rate of melting, and perhaps that of freezing also, is determined by the thickness and the thermal conductivity of such blankets, rather than by any inherent slowness with which the substance can change from one phase to the other (cf. p. 624).

In this connection, the excitement created in 1880-1882 by T. Carnelley's extravagant claim to have heated ice above 100 °C may be of interest. 114

¹¹¹ Hawkes, L., Nature, 123, 244 (1929).

¹¹³ Fisher, E. A., J. Phys'l Chem., 28, 360-367 (1924).

¹¹⁸ Barnes, H. T., "Ice Engineering," p. 2-3, 1928.

118 Barnes, H. T., "Ice Engineering," p. 2-3, 1928.

114 See Carnelley, T., Nature, 22, 434-435, 510-511 (1880); 23, 341-344 (1881); Chem. News, 42, 130, 313 (1880); Proc. Roy. Soc. London, 31, 284-291 (1880-81); Ber. deut. chem. Ges., 13, 2406-2407 (1880); J. Chem. Soc., 41, 317-323 (1882). Pettersson, O., Ber. deut. chem. Ges., 13, 2414-2144 (1880); Nature, 24, 167-169 (1881). Meyer, L., Ber. deut. chem. Ges., 13, 1831-1833 (1880); 14, 718-722 (1881). Wüllner, A., Ann. d. Physik (Wied.), 13, 105-110 (1881). And many other articles in Nature, vols. 22, 23, and 24. Certain of the articles are entitled "Hot Ice."

Rate of Freezing and of Melting.

The terms "rate of freezing," "quickness of freezing," "velocity of freezing," and their equivalents are essentially vague and indefinite when they stand alone, and have been used in different senses at different times. They may be used (1) to cover the rate of thickening of an ice sheet upon a pond, or (2) for the rate at which ice forms on an exposed surface of water, or (3) for the time that elapses between the exposure of water to chilling conditions and the initial appearance of ice in the water, or (4) for the rate at which crystallization proceeds along a narrow column of supercooled water; and they have been used in all these ways.

- 1. The rate of thickening of an ice sheet has been considered in Section 59 (p. 407).
- 2. In an exposition of his trihydrol theory of ice formation, H. T. Barnes 115 refers to certain experiments in which he periodically removed and measured the amount of ice that had formed on the surface of a tank of water since the preceding removal. He found that the rate of freezing, as so measured, decreased from one period to the next, finally becoming zero. If the water were heated to room temperature and then cooled again, ice would form as before. He regarded these observations as indicating that "water may be exhausted of its ice-forming power," "that a nucleus is required for the colloidal ice mass, and after exhausting these nuclei, the formation of ice is rendered difficult," and that time is "required for the restoration of the trihydrol in solution and at the temperature of freezing it is considerably slower than at higher temperatures." Except for a figure and its legend, given by T. C. Barnes and T. L. Jahn, 116 the details of these experiments seem to have remained unpublished. That is most unfortunate. The little that has been published is quite unconvincing and suggests that due attention was not given to important details. For example, the amount of ice that will form in a given time, once freezing is initiated, will be greatly affected by the heat capacity and initial temperature of the tank and its contents, and by the rate at which heat is abstracted from them by the chilling arrangement. But we are given no information that will enable one to form an estimate of these quantities, or of their variations from time to time; and there is no suggestion that they need to be considered. Moreover, the statement regarding the slow restoration of the trihydrol cannot be accepted without much better evidence than has been given us: in fact, it is most improbable.
- 3. T. C. Barnes and T. L. Jahn ^{116, 117} reported that, under the same conditions and starting from the same temperature, water from freshly melted ice freezes more quickly than that from freshly condensed steam. And they interpret this as confirming the conclusion of H. T. Barnes ¹¹⁵ that near and below 0 °C the recovery of equilibrium between the several polymers of H₂O is slow. Here again information regarding experimental

¹¹⁵ Barnes, H. T., Scientific Monthly, 29, 289-297 (1929).

¹¹⁶ Barnes, T. C., and Jahn, T. L., Quart. Rev. Biol., 9, 292-341 (1934).

¹¹⁷ Barnes, T. C., and Jahn, T. L., Proc. Nat. Acad. Sci., 19, 638-640 (1933).

details is meager, and satisfactory checks are wanting. There is no indication that any attention has been paid to the great variations that have been found in the extent to which water can be supercooled; variations that force one to suspect, at least, that the extent of possible supercooling truly

Table 282.—Velocity of Crystallization of Supercooled Water

By the velocity of crystallization, we here mean the linear velocity, v, with which freezing, initiated at one end, proceeds along the length of a glass tube filled with supercooled water and continuously immersed in a bath at the same temperature, t, as the water at the instant that freezing began. The internal diameter of the tube is d, the wall thickness is w.

Unit of v = 1 mm/sec; of d and w = 1 mm. Temp. = t °C

	One or v	Walton	and Judd.a	Tempi V O	
$d\rightarrow$,			2	E
$w \rightarrow$	7 2.5	i	1 .5	3. 3.	25
-t	v	-t	v	- t	v
2.00	5.27				
3.61	8.07				
4.67	11.9				
5.86	17.8				
6.18	19.1	6.17	39.2	6.10	24.0
7.10	44.4			6.60	25.2
7.50	51.3	7.65	63.0	7.58	32.2
8.19 8.38	69.2				
8.38	85.5			8.58	39.1
9.07	114.0	9.92	91.6		
Tu	mlirzª	Harts	mann ^a	TH	3°
$d \rightarrow$	18		:0 3	1	.2 .8
$v \rightarrow$.8
-t	\boldsymbol{v}	-t	T)	-t	\boldsymbol{v}
0.74	0.37	0.5	2.3		
1.12	1.44	0.8	3.3		
1.40	2.20	1.0	4.0		
1.54	2.76	1.5 1.9	6.0 8.0		
1.62	2.92				
2.00	3.32	1.9	8 5 9.7		
2.40	4.49	2.0 2.0	9.7 9.7		
2.54 2.67	5.24 5.58	3.5	20.0	3.2	11.18
2.71	5.77	3.5	20.3	4.2	16.2
2.90	7.06	5.0	29.2		23.8
3.20	7.00 7.47	5.0	20.2	6.2	30.5
3.49	10.23	7.0	46.2	7.2	41.3
3.64	11.28	7.0	46.7	83	52.0
4.14	16.93	7.0	10.7	5.2 6.2 7.2 8.3 9.3	55.0
4.20	18.15			10.3	61.2
4.60	22.07			11.3	70.3
₩.00	22.07			12.3	84.0
				13.4	96.8

* References:

Hartmann, R., Z. anorg. allgem. Chem., 88, 128-132 (1914).
TB Tammann, G., and Büchner, A., Idem, 222, 12-16 (1935).
Tumlirz, O., Sitz. Akad. Wiss Wien (Abt. IIa), 103, 266-276 (1894). As quoted by Walton, J. II., and Judd, R. C., J. Phys'l Chem., 18, 722-728 (1914).

 $^{^{}b}$ For temperatures above -6 $^{\circ}$ C they used tubes 3 mm in internal diameter; the wall thickness was not found reported.

varies from specimen to specimen, depending upon some inclusion foreign to the water itself, and not merely upon the thermal treatment of the water. See remarks in the note at the beginning of this Section, p. 638.

4. As the rate of melting of ice immersed in water depends upon the rate at which heat is delivered to the ice, so the linear velocity (see Table 282) with which freezing proceeds along the length of a tube filled with supercooled water and immersed in a bath at a constant temperature measures the rate at which heat is removed, rather than a characteristic property of the water-substance. But it is possible that, with the available facilities, heat cannot be removed from ice at a rate that is greater than some fixed amount determined by the characteristics of water. In that case the linear velocity of the freezing will approach a maximum as the rate of abstraction of heat is increased, and that maximum will be determined by some property, or group of properties, of water. The recorded observations give no indication of such a limiting value.

For a discussion of the linear velocity of crystallization, see the references given in Table 282 and the recent papers by R. Kaischew and I. N. Stranski ¹¹⁸ and T. Förster. ¹¹⁹

Rate of Melting: Effect of Tension.—The effect of tensile stress upon the rate at which ice melts when exposed to air slightly above 0 °C has been studied by O. Fabian. Using cylinders of ice, all of the same size (diameter = 5.4 cm) but differently loaded, and all exposed simultaneously in a room in which the air temperature was 0.9 °C, he found as follows:

Load	0	25	50	kg*
Loss in weight	20	17.5	13	g/hr

Crystalloluminescence.

Statements ¹²¹ to the effect that Pontus, in 1833, observed that water luminesces when it freezes—that it exhibits crystalloluminescence—appear to be incorrect.

The announcement of Pontus's observation ¹²² states that when a glass bulb with a small tubular neck 1 or 2 cm long is completely filled with water, wrapped with cotton soaked with ether, and placed in a receiver which is then exhausted, then a spark, visible in full daylight, jumps from the neck some moments before freezing occurs ("quelques instans avant la congélation une étincelle bien visible en plein jour s'échappe du petit tube qui termine l'ampoule"). This is certainly not the description of crystallo-luminescence; and that no such brilliant light accompanies either the freezing of water or the formation of frost, however rapid the process, is to be inferred from the total absence of any mention of it in the voluminous records treating of those processes. Any one can readily satisfy himself

¹¹⁸ Kaischew, R., and Stranski, I. N., Z. Physik. Chem. (A), 170, 295-299 (1934).

¹¹⁹ Förster, T., Idem, 175, 177-186 (1936).

¹²⁰ Fabian, O., Report. Exper. Physik (Carl), 12, 397-404 (1876).

¹²¹ Trautz, M. Z. physik. Chem., 53, 1-111 (1905). Mellor, J. W., "A Comprehensive Treatise on Inorganic and Theoretical Chemistry, vol. 1, p. 465, London, Longmans & Co., 1922.

122 Pontus, J. chim. med., 9, 429-430 (1833) → Ann. d. Physik (Pogg.), 28, 637 (1833).

that if there is any crystalloluminescence involved in such freezing it is certainly very weak and can in no sense be confounded with the phenomenon observed by Pontus. His entire description indicates that the spark observed was a secondary phenomenon.

98. Transition of Ice to Ice

(See also, Types of Ice, Section 57.)

The numerical data of various kinds pertaining to the transition of ice to ice having been given elsewhere (external work and change in volume and latent heat, Section 95; transition temperature, Table 270; phase diagram, Section 93), only a few descriptive items remain for this section.

Ice-I can be carried into the domains in which ice-II and ice-III are stable, but ice-II cannot be carried into that of either ice-III or ice-V; ice-III can be carried into the domains of ice-I, ice-II, and ice-V; ice-IV occurs between the domains of ice-III and ice-VI, within that in which ice-V is stable, but it is totally unstable with reference to ice-V, vanishing entirely if ice-V appears (B600)*; ice-V can be carried into the domains of ice-II and ice-III, and into that of ice-VI at temperatures well below that of the triple point, but near that point it cannot be carried the slightest distance into the domain of ice-VI; and ice-VI can be carried far into that of ice-V, and can be kept there for a considerable time without changing into ice-V.

The velocity with which one type of ice changes to another, when carried over into the pressure-temperature domain in which the second is the stable form, ranges from explosive rapidity near the triple point of higher temperature to extreme sluggishness at lower temperatures. For example, when ice-III is formed from ice-I at temperatures above $-30\,^{\circ}\text{C}$, the reaction runs "with explosive velocity," sometimes producing a sharp, audible click (B478); but at $-70\,^{\circ}\text{C}$ the change is so slow as to be not appreciable within 4 hours, even though the pressure be several hundreds of kg*/cm² from the equilibrium one (B476). Since the latent heat varies but little with the temperature, it is obvious that something else is of prime importance in the regulation of the speed of transition. What it is, is not known (B535).

The possibility of carrying ice-VI into the domain of ice-V seems to depend upon conditions that one might expect to be quite unessential. For example: At temperatures above -25 °C, but below the melting line, ice-VI could regularly be carried far into the domain of ice-V, but if bits of Jena glass were placed in the water, then ice-V promptly appeared (B503-506). At lower temperatures ice-V could be obtained from either ice-II or ice-III without the presence of glass (B506).

For further details, see the papers on which these statements rest-

^{*} For such references, see end of this section.

P. W. Bridgman.¹²³ Such references as (B593-506), occurring in the text, refer to the pages of these papers. All but one (B600) refer to the first; that one, obviously, to the second.

99. MISCELLANEOUS CHANGES ACCOMPANYING PHASE TRANSITION

Here are assembled those changes accompanying phase transition that do not fit satisfactorily into the preceding sections, and in general, only those changes that have been directly observed. Others will be found recorded in the pertinent sections, or may be derived from data given therein.

Table 283.—Change in Refraction with Change in Phase

From the observations of others, P. Hölemann 124 has computed the following values of the molecular refraction: $R \equiv M(n^2-1)/\rho(n^2+2)$, where M = molecular weight (18.0154), n = index of refraction, and $\rho =$ density. Subscripts v and l indicate that the value refers to the vapor and to the liquid, respectively.

Unit o	$f \lambda = 1 m\mu = 10^{-7} cm$; of $R = 1 \text{ cm}^3/3$	g-mole
λ	R_v	Ri	$100 \ (R_{l}-R_{v})$
435.8	3.8262	3.7851	4.11
467.8	3.8041	3.7637	4.04
480.0	3.7969	3.7571	3.98
501.6	3.7854	3.7463	3.91
508.6	3.7820	3.7427	3.93
546.1	3.7660	3.7269	3.91
587.6	3.7510	3.7124	3.95
643.9	3.7371	3.6965	4.06
656.3	3.7344	3.6931	4.13
667.8	3.7321	3.6903	4.18

Table 284.—Change in Absorption Spectrum with Change in Phase

The bands in the absorption spectrum of water vapor are more numerous than those in the spectrum of either water or ice, and have a very complicated "fine structure"; those in the spectrum of water and of ice are nearly devoid of "fine structure," and those of water may have sharp edges whereas those of ice do not. The wave-length corresponding to the maximum of absorption in a given band common to all three phases increases as the substance passes from vapor to liquid to solid. (cf. MRB.^a) This increase in λ on passing from one phase to the next is here called $\Delta\lambda$.

For ice there is a band at $\lambda = 4.75 \,\mu$, for water one at 4.7, but there is no corresponding band for water-vapor (E1^a).

For the vapor the bands at $\lambda = 1.44$ and $2.00 \,\mu$ are stronger than for water, but the reverse is true of those at $\lambda = 0.97$ and $1.20 \,\mu$ (Co^a).

Bridgman, P. W., Proc. Am. Acad. Arts Sci., 47, 441-558 (1912); J. Phys'l Chem., 3, 597-605 (1935); Idem, 5, 964-966 (1937).
 Hölemann, P., Z. physik. Chem. (B), 32, 353-368 (1936).

Table 284.—(Continued)

Unit of λ and $\Delta\lambda = 1$ $\mu = 10^4$ A = 10-4cm

Vapor 100 °C	Water	Δλ	Refa	220 °C Vaj	20 °C λ	97 °C Wa	ter 20 °C	Ref*
1.404 1.885 2.661	1.475 1.970 2.916	0.071 0.085 0.225	Dr Dr Pa	1.414 1.881 2.600	1.414 1.881 2.620	1.468 1.945	1.475 1.954 2.950	St St St
w	ater	Ord λ (0 °C) —	Ice Ext	Or	d Δλ	Ext	Refa	
0. 0.	745 845 98 215	0.79 0.89 1.02 1.26	0.81 0.92 1.06 1.29	0.0)45)45)4)45	0.065 0.075 0.08 0.075	Pi Pi Pi Pi	

* References:

- Collins, J R., Phys. Rev. (2), 20, 486-498 (1922). Co
- Dreisch, T., Z. Physik, 30, 200-216 (1924).

 El Ellis, J. W., Phil. Mag. (7), 3, 618-621 (1927).

 MRB McLennan, J. C., Ruedy, R., and Burton, A. C., Proc. Roy. Soc. London (A), 120, 296-302 (1928).
- Paschen, F., quoted by Dr, presumably from Ann. d. Physik (Wied.), 53, 334-336 Pa
- Plyler, E. K., J. Opt. Soc. Amer., 9, 545-555 (1924). Stansfeld, B., Z. Physik, 74, 460-465 (1932). Pl
- St

Raman Spectra.

The displacements of the most prominent lines or bands as the phase and temperature are changed are shown in Table 285. I. R. Rao 125 regards these changes as arising from changing proportions of the molecules (H₂O), (II₂O)₂, and (H₂O)₃ present in the substance. To (H₂O)₈ he ascribes $\lambda_R = 3.13 \,\mu \, (\delta \nu = 3195 \, \text{cm}^{-1})$; to $(H_2O)_2$, $\lambda_R = 2.93 \,\mu \, (\delta \nu = 1.00 \, \text{m}^{-1})$ 3413 cm⁻¹); and to (\dot{H}_2O), $\lambda_R = 2.77 \,\mu$ ($\delta_V = 3610 \,\text{cm}^{-1}$). On the other hand, G. B. B. M. Sutherland 126 has attempted to explain the changes in terms of but two types of molecules-H2O and (H2O)2-, ascribing the $\delta v = 3200 \text{ cm}^{-1} \text{ line to } (H_2O)_2.$

In the spectrum of the vapor there is only a single prominent line $(\delta v = 3655 \text{ cm}^{-1})$, which is fine and sharp; in that of water, there are no prominent sharp lines, but there are two prominent bands, one very broad and complex; in that of ice near 0 °C, there are two bands, each much narrower than the water bands, and at about -190 °C there are two fairly sharp lines, one intense.127

Water of crystallization gives a Raman spectrum that is very similar to that of ice, but the lines are sharper. 128

¹²⁵ Rao, I. R., Proc. Roy. Soc. London (A), 145, 489-508 (1934).

¹⁹⁸ Sutherland, G. B. B. M., Idem, 141, 535-549 (1933).

¹⁷⁸ Bhagavantam, S., Indian J. Phys., 5, 49-57 (1930). Cabannes, J., and de Riols, J., Compt. rend., 198, 30-32 (1934). Daure, P., and Kastler, A., Idem, 192, 1721-1723 (1931). Ganesan, A. S., and Venkateswaran, S., Indian J. Phys., 4, 195-280 (1929). Rao, I. R., Idem, 3, 123-129 (1928). Nature, 225, 600 (1930); Proc. Roy. Soc. London (A). 145, 489-508 (1934); Phil. Mag. (7), 17, 1113-1134 (1934). Kohlrausch, K. W. F., "Der Smekal-Raman Effekt," Springer, Berlin, 1931. ¹²⁸ Cabannes, J., and de Riols, J., ¹²⁷ Ganesan, A. S., and Venkateswaran, S., ¹²⁸ Krishnan, K. S., Indian J. Phys., 4, 131-138 (1929); Kohlrausch, K. W. F., ¹²⁷ and many others.

Table 285.—Change in Raman Spectrum with Change in Phase

•		Unit of	$\delta v = 1$	cm-1; of	$\lambda_R = 1 \mu$	= 10-4	cm			
Phase			δν					λ <i>R</i>		
I. I. R. R:	ao. ¹²⁵									
Ice		3196	3321				3.13	3.01		
Water 0 °C			3321	3502				3.01	2.86	
Water 98 °C				3 466					2.88	
Water-vapor					3655°					2.74°
II. A. S.	Ganesan	and S.	Venka	iteswara	ın. ¹²⁷					
Ice		3193	3391	3549	5349		3.13	2.95	2.82	1.85
Water	2355	3199	3453	3609	5502	4.25	3.13	2.90	2.77	1.82
III. Sever	al obser	vers.								
Water		3214	3440	3604			3.11	2.91	2.77	
Water-vapor					3655					2.74
^a Measurement by P. Daure and A. Kastler. ¹²⁷										

Magnetic Susceptibility.

At the request of Piccard, G. Foex measured relatively the specific susceptibility (χ) for the same specimen of water when frozen and when liquid. He found that at the moment of freezing the numerical value of χ decreased by 2.4 per cent of its value for the liquid.¹²⁹ T. Ishiwara ¹³⁰ reported a minute change in the same direction, and of approximately the same magnitude, and more recently, B, Cabrera and H. Fahlenbrach ¹³¹ found 2.2 per cent.

¹²⁰ Piccard, A., Arch. Sci. phys. ct nat. (4), 35, 209-231, 340-359, 458-482 (1913).

¹⁸⁰ Ishiwara, T., Sci. Rep. Tohoku Univ., Sendai (1), 3, 303-319 (1914).

¹⁸¹ Cabrera, B., and Fahlenbrach, H., An. Soc. Esp. Fis. y Quim., 31, 401-411 (1933).

V. Miscellanea

100. MISCELLANEOUS PHENOMENA AND DATA

Penetration of Solids by Water.

If either glass or quartz is exposed for 5 or 10 minutes to water (or to certain other liquids) at a pressure of 15 000 atmospheres, or over, and the pressure is then suddenly released, the glass or quartz will be broken, perhaps shattered. This is explained by the gradual penetration of the compressed liquid into the solid, which is unable to withstand the resulting stress when the outer pressure is removed.1

Thermal Anomalies of Water.

- M. Magat 2 has concluded that the existing data indicate that most of the physical properties of water exhibit thermal anomalies in the neighborhood of 35 to 40 °C.
- A. P. Wills and G. F. Boeker 3 and S. Seely 4 report that the thermal variation of the magnetic susceptibility of water is anomalous in the range 35 to 55 °C, the anomaly being marked at each extreme.
- G. Tammann 5 reports that the existing data show that each of many properties of water has either a maximum or a minimum value near 50 °C.
- J. Timmermans and H. Bodson 6 find that their determinations of the surface tension of water show a clear anomaly at 13 °C.

Impact of Solids upon Water.

The resistance offered by water to the impact upon its surface of solids of various forms has been studied by S. Watanabe.⁷

Table 286.—Volume of the Water Meniscus: Special

(See also Table 287.)

By the volume of the water meniscus is meant the volume (v) of water that lies above the horizontal plane that is tangent to the bottom of the meniscus in a vertical, cylindrical tube of circular cross-section (radius = r). the angle of contact being zero. The quantity $l \equiv v/\pi r^2$ may be called the

- 1 Poulter, T. C., and Wilson, R. O., Phys. Rev. (2), 40, 877-880 (1932).
- ² Magat, M., Jour. de Phys. (7), 6, 179-181 (1935); 6, 64S-65S (1935); Trans. Faraday Soc., 33, 114-120 (1937). Especially the first.
 - 8 Wills, A. P., and Boeker, G. F., Phys. Rev. (2), 46, 907-909 (1934).
 - 4 Seely, S., Idem, 52, 662 (L) (1937).
 - ⁵ Tammann, G., Z. anorg. allgem. Chem., 235, 49-61 (1937).
 - 6 Timmermans, J., and Bodson, H., Compt. rend., 204, 1804-1807 (1937).
- ⁷ Watanabe, S., Sci. Papers Inst. Phys. Chem. Res. (Tokyo), 23, 118-137, 202-209, 249-255 (1934).

Table 286.—(Continued)

equivalent height of the meniscus; the values of l commonly given for water in tables of constants are those (l_B) determined experimentally by Robert Bunsen, and published in his "Gasometrische Methoden," 1877 8; they are all too small. The following values of l and v have been computed in part from the table given by F. A. Gould 9 and in part from that published by S. Sugden. Oraphical interpolation and smoothing were employed.

In the experimental work of W. Bein 11 the angle of contact was not zero.

 $a^2 \equiv 2\gamma/(\rho - \sigma)\dot{g} = 0.15$ cm² at 15 °C, 0.14 cm² at 50 °C (see Table 225); $\gamma =$ surface tension. If 2r < 1 mm, l is essentially equal to r/3; if 2r > 5 cm, then, within less than 0.5 per cent, v/r = 0.438 cm² when $a^2 = 0.14$ cm², and 0.470 cm² when $a^2 = 0.15$ cm².

Unit of	$a^2 = 1 \text{ mm}^2 = 0$.01 cm ² ; of r	and of $l=1$	mm; of $v = 1$	mm ³ = 0.001 cm ³
$a^2 \rightarrow 2r$		15	l_B	14	- 7'
	0.16 0.32 0.47	0.16 0.32 0.47	12	0.13 1.04 3.3	0.13 1.04 3.3
4 5	0.60 0.72	0.61 0.73		7.5 14.1	7.7 14.3
1 2 3 4 5 6 7 8 9	0.82 0.90 0.97 1.00 1.06	0.83 0.92 0.99 1.00 1.07		23.2 34.6 48.8 64 83	23.5 35.4 49.8 64 84
11 12 13 14 15	1.10 1.12 1.13 1.14 1.12	1.12 1.15 1.17 1.18 1.16	1.10 1.03	104 127 150 175 199	106 130 155 181 206
16 17 18 19 20	1.13 1.12 1.12 1.11 1.10	1.17 1.16 1.16 1.15 1.14	0.97 0.91 0.87 0.84 0.82	226 254 284 316 346	234 264 295 326
22 24 26 28 30	1.08 1.03 0.98 0.94 0.89	1.12 1.09 1.04 0.99 0.94	0.02	409 467 523 575 628	358 426 493 552 610 665
32 34 36 38 40	0.84 0.80 0.76 0.73 0.69	0.90 0.86 0.82 0.78 0.74		678 729 779 825 870	722 779 833 885 936
42 44 46 48 50	0.66 0.63 0.60 0.58 0.56	0.71 0.68 0.65 0.62 0.60		916 960 1004 1048 1092	985 1032 1080 1127 1174

⁸ Bunsen, Robert, "Gesammelte Abhand.," vol. 2, p. 364, Leipzig, 1904.

⁹ Gould, F. A., Int. Crit. Tables, 1, 73 (1926).

¹⁰ Sugden, S., J. Chem. Soc. (London), 119, 1483-1492 (1921).

¹¹ Bein, W., Z. Inst.-kunde, 48, 161-163 (1928).

Table 287.—Volume of the Water Meniscus: General (Adapted from A. W. Porter.¹² See also Table 286.)

These values for vertical circular cylinders and for cones diverging upwards are based upon the tables of Bashforth and Adams ¹³ if $r/\beta < 5$, and upon the formula derived by the late Lord Rayleigh 14 for large tubes if $r/\beta > 5$. They assume that the contact angle is zero, otherwise they apply to any fluid, an appropriate value being assigned to β . Here, $\beta^2 \equiv \gamma$ $(\rho - \sigma)g = a^2/2$, a^2 having the same significance as in Table 286; $\gamma = \text{sur-}$ face tension; $2\phi = \text{vertex}$ angle of the cone; r = radius of the tube where met by the meniscus; h = the elevation of that intersection above the (horizontal) plane that is tangent to bottom of the meniscus; v = volumeof liquid lying above that plane; v_a = value of v as obtained from Table 286.

	Unit of \$	and $a = 1$ mm; o	f v = 0.001 cm	$n^3 = 1 \text{ mm}^3$	
 Cylinders. 					
2 β ² (- c		1	4 ———		15
r/B	$v/\pi \beta^3$	v	Va	ข	Va
0.8853	0.2053	11.9	12	13.2	13
1.8687	1.4092	82.0	81	90.9	89
2.1688	1.9857	115.5	115	128.1	127
2.4074	2.4966	145.3	144	161.1	160
2.9192	3.6847	214.4	211	237.8	234
3.1646	4.3264	251.7	247	279.2	274
	9.077	528.1	535	585.7	592
6	11.4635	667.0	672	739.7	747
ž	13.7109	797.7	803	884.7	894
5 6 7 8	15.8528	922.4	923	1022.9	1028
9	17.928	1043.1	1040	1156.8	1157
10	19.966	1161.7	1156	1288.4	1286
II. Cones.					
III Concis.	$-\phi = 30^{\circ}$			\(\phi = 45^\circ	
+/β	h/β	$v/\pi\beta^3$	τ/β	h/β	$v/\pi\beta^3$
0.7916	0.4292	0.0425	0.6658	0.2660	0.0067
1.3507	0.6617	0.2053	1.1780	0.4388	0.0796
1.745	0.7841	0.4272	1.5566	0.5414	0.1826
2.0428	0.8563	0.6570	1.8484	0.6059	0.2847
2,2804	0.9031	0.8773	2.088°	0.6492	0.4234
3.0372	1.0025	1.7650	2,4464	0.7030	0.6563
0.0072	2.0020	217 0.00	2.5923	0.7206	0.7644
			2.8376	0.7456	0.9080
^a This appears	once as 2.0	088 and again as	2.0833.		

Table 288.—Radiation from an Ideal Black-body Radiator (Adapted from F. E. Fowler.14a)

The energy radiated per unit time by a unit area of an ideal (blackbody) radiator at $t \, {}^{\circ}$ C is $10^{n}R$. If there is present another body not at

¹² Porter, A. W., Phil. Mag. (7), 14, 694-700 (1932). The study is continued and extended in Porter, A. W., Trans. Faraday Soc., 29, 702-707, 1307-1309 (1933); Phil. Mag. (7), 17, 511-517 (1934).

¹⁸ Bashforth and Adams, "An Attempt to Test the Theory of Capillary Action," Cambridge Univ. Press, 1883.

¹⁴ Rayleigh, Lord, Proc. Roy. Soc. London (A), 92, 184-195 (1916). ^{24a} Fowle, F. E., "Smithsonian Physical Tables," 8th revised edition, p. 313, Table 307, Washington, 1933.

Table 288.—(Continued)

absolute zero (-273 °C), then radiation will be received from it, and the net loss of energy by the ideal radiator will be correspondingly less than R. $10^nR = \sigma T^4$, where $\sigma = 5.73 \ 10^{-5} \text{erg/cm}^2 \text{sec}$, T = (273 + t) °K, temperature = t °C; 1 kcal = 1000 cal = 4185 joules = 4185·10⁷ ergs.

Unit→	1 erg/c	m²sec	1 cal/c	cm²sec	1 kcal/c	m²hr
ŧ	R	21	R	n	R	n
0	3.19	5	7.66	-3	2.76	2
100	1.11	6	2.66	-2	9.58	2
200	2.87	6	6.89	-2	2.48	3
300	6.18	6	1.48	-1	5.33	3
400	1.18	7	2.83	-1	1.02	4
500	2.05	7	4.92	-1	1.77	4
600	3.33	7	7.99	-1	2.88	4
700	5.14	7	1.23	0	4.43	4
800	7.60	7	1.82	0	6.55	4
900	1.11	8	2.66	0	9.58	4
1000	1.50	8	3 60	0	1.30	5
1100	2.04	8	4.87	0	1.75	5
1200	2.70	8	6.45	0	2.32	5
1300	3.51	8	8.39	Ö	3.02	5
1400	4.49	8	1.07	ĭ	3 86	5
1500	5.66	8	1.36	1	4.90	5
1600	7.05	8	1.69	1	6.07	5
1700	8.68	8	2.08	1	7.47	5
1800	1.06	ğ	2.53	1	9.11	5
1900	1.28	9	3 06	1	1.10	6
2000	1.53	9	3.67	1	1.32	6

Vision under Water.

R. E. Cornish ¹⁵ has described a spectacle lens that facilitates vision by an eye immersed in water. Such immersion greatly reduces the refraction of the eye, the index of refraction of the aqueous humour being nearly that of water.

Sea-water.

Properties of sea-water that are analogous to those of pure water will be found in the appropriate sections for water, if given at all in this compilation. Data pertaining to its composition and temperature are given here.

The composition of sea-water varies from place to place, and from time to time, depending upon the evaporation and the inflow of fresher water from streams, ice-bergs, and precipitation. G. Wüst ¹⁶ has stated that the salinity (S) of the surface layer of the sea far from shore is given by the formula S=35.74+0.0126(E-P), where E and P are, respectively, the rate of evaporation and of precipitation (unit of S=1 g salt per kg of sea-water; of E and P=1 mm per day). For values of E, see Table 277. Sea-water averages about 35 g of salts per kg; all chemical elements are

¹⁸ Cornish, R. E., J. Opt. Soc. Amer., 23, 430 (1933).

¹⁶ Wüst, G., Metcor. Z., 38, 188-190 (1921) ← Veröffentl. Inst. Meereskunde (N. F.), geog.-naturw. Reihe, Heft 6, (1920).

represented, most of them by very minute amounts. The relative amounts of the more abundant are shown in Table 289.

In the surface layers of the oceans the salinity increases from about 35.1 at the equator to about 36 near latitude 25°, and then decreases, reaching about 30 near 70° N and 33 near 70° S; in general the salinity is somewhat greater in the southern hemisphere than at the corresponding latitude in the northern one. Our knowledge of the distribution of salt throughout the depth of the oceans is very imperfect.¹⁷

The mean temperature of the surface layers of the oceans is about 27 °C at the equator, 20 °C at latitude 30°, 4.8 °C at 60° N, 0.0 °C at 60° S, and -1.7 °C at 80° N and 80° S; in general the temperature is lower in the southern hemisphere than in the northern. The mean temperature of the oceans from top to bottom is about 4.8 °C at the equator, 3 °C at latitude 45°, -0.6 °C at 75° N, and +0.9 °C at 75° S.¹⁷

Table 289.—Composition of the Salt of Sea-water 18

Cl = total mass of chlorine per unit mass of sea-water; m_s and Cl_s = mass of the indicated salt and of the associated chlorine, respectively, per unit mass of sea-water; S = salinity = total m_s . From the values tabulated, it follows that S = 34.4 g/kg, Cl = 18.99 g/kg, S = 1.812Cl, Cl = 0.552S. The actual value of S is subject to variations, but the ratios m_s/S , and S/Cl are essentially constant.

Unit of S,	Cl, and ms	== 1 g/kg	
Salt	972 -	m = /S	Cl.
NaCl	26.9	0.783	16.33
MgCl ₂	3.2	0.094	2.38
MgSO ₄		0.064	
CaSO,	1.4	0.039	
KC1	0.6	0.017	0.28
Rest	0.1	0.003	
Total	34.4	1.000	18.99

Surprises.

Here are listed a few of the things referring to water that seem to the compiler to be thought-provoking. They are merely those he happened to jot down. They, and probably others equally worthy of a place in this list, have been considered elsewhere in connection with related phenomena.

- 1. The density of water that has stood in contact with carbon or with thoria is abnormal ¹⁹ (p. 225).
- 2. The vapor pressure of water in contact with catalysts is abnormally great. Prolonged heating, followed by a return to the initial temperature, affects that increase, and weeks may be required for it to return to its pristine value.²⁰ The existence of this effect has been questioned (p. 560).

¹⁷ Krümmel, O., "Handb. d. Ozeanog.," Bd. 1, 1907.

¹⁸ Krümmel, O., and Ruppin, E., Wiss. Mecresunters. (N. F.), 9, (Abt. Kiel), 27-36 (1906).

¹⁹ Peel, Robinson, and Smith, Nature, 120, 514-515 (1927).

²⁰ Baker, H. B., J. Chem. Soc. (London), 1927, 949-958 (1927).

- 3. Under the same conditions as in the preceding, the degree of association of the water, as computed by the method of Ramsay and Shields from the temperature coefficient of the surface tension, is abnormally great, and shows similar variations after prolonged heating.²⁰ The existence of this effect has been questioned (p. 172).
- 4. Different samples of nominally identical water may have different indices of refraction ²¹ (p. 279).
- 5. The density of ice-I seems to be subject to well-marked variations (p. 463).

Interpolation.

If the values f_0 , f_1 , f_2 , f_3 ... of f(x) corresponding respectively to the values x_0 , x_1 , x_2 , x_3 ... are known, the x's progressing by equal steps $(x_1 = x_0 + s, x_2 = x_1 + s, x_3 = x_2 + s, ...)$, then the value of $f(x_n + h)$, $x_n + h$ lying between x_n and $x_n + s$, may be found by means of formula (1)

$$f(x_n + h) = f_n + ka_n + \frac{k(k-1)}{2!}b_n + \frac{k(k-1)(k-2)}{3!}c_n + \frac{k(k-1)(k-2)(k-3)}{4!}d_n + \dots$$

$$= f_n + k\left\{a_n - \frac{1-k}{2}b_n + \frac{(1-k)(2-k)}{6}c_n - \frac{(1-k)(2-k)(3-k)}{24}d_n + \dots\right\}$$
(1)

in which the symbol! (read "factorial") indicates that the continued product of all the integers from one to that appearing before the symbol is to be taken $(4! = 1 \times 2 \times 3 \times 4 = 24)$; k = h/s, a quantity that is less than unity; and a_n , b_n , c_n , d_n , ... are the successive tabular differences (Δ) associated with x_n , as given by the following scheme, in which $a_0 = f_1 - f_0$. $a_1 = f_2 - f_1 \dots b_0 = a_1 - a_0$, $b_1 = a_2 - a_1$, ... $c_0 = b_1 - b_0$, $c_1 = b_2 - b_1$, ..., etc. The subtraction must always be made in the direction here indicated, and the proper sign must be given to the difference.

x	f(x)	Δ_{t}	Δ_2	Δ_3	Δ_1	Δ_5	etc.
x_0	fo	a_0	b_{o}	Co	d_{0}		
x_1	f_1	a_1	b_1	c_1	d_1		
x_2	f <u>.</u>	a_2	b_2	C2	d_2		
x_3	f_{a}	a_{a}	$b_{\mathbf{a}}$	Ca	d_3		
	•						
•	;	•	.•	•	:	•	
x_n	†n	a_n	b_n	Cn	d_n	•	
•	•	•	•	•	•	•	
•	•	•	•	•	•	•	
•	•	•	•	•	•	•	

²¹ Damien, B. C., Ann. Sci. École Norm. Sup. (2), 10, 233-304 (272-278) (1881) → Jour. de Phys. (1), 10, 198-202 (1881). v. d. Willigen, V. S. M., Arch. Mus. Teyler, 1, 74-116, 161-200, 232-238 (1868).

Index

This is a subject index. That is, each page reference following an entry indicates the location of information that bears in some manner on the subject covered by the entry, no matter what may be the actual wording of the text. In many cases the index entry is a more comprehensive term than the pertinent one in the portion of the text to which reference is made.

In order to facilitate the use of the text, the same information has, in many cases, been indexed under several entries, each being in some manner appropriate. Nevertheless, the user will now and again fail to find an entry that he expects. In such cases the information sought may be found under a synonym of that entry, or under some more comprehensive term; if not so found, the section, or sections, in which it should be given may be found by reference to the Table of Contents. If this also fails, the inference is that the compilation does not contain the information sought.

A main entry that is followed by subordinate (indented) ones carries a page reference only if it refers to a main section of the compilation or if there are pertinent items not specifically covered by any of the subordinate entries, and only for such items. Such partial references are enclosed by parentheses.

Symbols used to denote mathematical and physical quantities and units of measure are not indexed individually. Those not explained in the text where used should be sought under the word "symbols."

Explanatory words indicating the nature of the quantity or of the information given, the field concerned, the phase of the substance, the independent variable (p, t, v, etc.) are enclosed in parentheses. Inversion of the natural order of the words is indicated in the usual manner, by capitalizing the first letter of what would naturally be the initial word, and placing a comma before it, thus: Association, Molecular = Molecular association.

The same index entry may cover many pages. If these are consecutive, then only the first page is given, unless there seems to be danger that the user may fail to notice over how many pages the pertinent information extends; in which case, either the extreme pages are indicated or the initial page is followed by a +. If the pages are not all consecutive, then the initial page, modified as just described, is given for each group of consecutive pages.

If a term is defined wherever used, its definition is, in general, not specifically indexed. In other cases, the page on which the definition occurs is indexed and followed by a d; if it is desired to indicate that other pertinent information is to be found on the same page the d is enclosed in

parentheses; otherwise it is not. If the same term is defined in each of several sections, then reference to each of these definitions is usually given, thus relieving the user of the necessity of turning to a distant portion of the volume.

The following abbreviations are used: cf. = compare; d = definition is on this page; eff. = effect of, or on; q.v. = which see = seek in the index the term preceding the q.v.; see + = see also; a + placed after a page number means "and following pages"; vapor = water-vapor.

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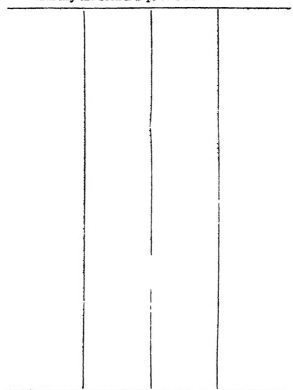
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